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(57) Abstract: The present invention relates to proteome chips comprising arrays having a large proportion of all proteins expressed in a single species. The invention also relates to methods for making proteome chips. The invention also relates to methods for using proteome chips to systematically assay all protein interactions in a species in a high-throughput manner. The present invention also relates to methods for making and purifying eukaryotic proteins in a high-density array format. The invention also relates to methods for making protein arrays by attaching double-tagged fusion proteins to a solid support. The invention also relates to a method for identifying whether a signal is positive.



GLOBAL ANALYSIS OF PROTEIN ACTIVITIES USING PROTEOME CHIPS

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RELATED APPLICATIONS

This application claims benefit of United States provisional application nos. 60/290,583, filed on May 11, 2001, and 60/308,149, filed on July 26, 2001, each of which is incorporated herein by reference in its entirety.

15 1. FIELD OF THE INVENTION

The present invention relates to proteome chips comprising arrays having a large proportion of all proteins expressed in a single species. The invention also relates to methods for making proteome chips. The invention also relates to methods for using proteome chips to systematically assay all protein interactions in a species in a high-throughput manner.

The present invention also relates to methods for making and purifying eukaryotic proteins in a high-density array format. The invention also relates to methods for making protein arrays by attaching double-tagged fusion proteins to a solid support. The invention also relates to a method for identifying whether a signal is positive.

2. BACKGROUND OF THE INVENTION

A daunting task in the post-genome sequencing era is to understand the functions, modifications, and regulation of every protein encoded by a genome (Fields et al., 1999, Proc Natl Acad Sci. 96:8825; Goffeau et al., 1996, Science 274:563). Currently, much effort is devoted toward studying gene, and hence protein, function by analyzing mRNA expression profiles, gene disruption phenotypes, two-hybrid interactions, and protein subcellular localization (Ross-Macdonald et al., 1999, Nature 402:413; DeRisi et al., 1997, Science 278:680; Winzeler et al., 1999, Science 285:901; Uetz et al., 2000, Nature 403:623;

Ito et al., 2000, Proc. Natl. Acad. Sci. U.S.A. 97:1143). Although these studies are useful, transcriptional profiles do not necessarily correlate well with cellular protein levels. Thus, the analysis of biochemical activities can provide information about protein function that complements genomic analyses to provide a more complete picture of the workings of a cell (Zhu et al., 2001, Curr. Opin. Chem. Biol. 5:40; Martzen, et al., 1999, Science 286:1153; Zhu et al., 2000, Nat. Genet. 26:283; MacBeath, 2000, Science 289:1760; Caveman, 2000, J. Cell Sci. 113:3543).

Several groups have recently described microarray formats for the screening of protein activities (Zhu et al., 2000, Nat. Genet. 26:283; MacBeath et al., 2000, Science 289:1763; Arenkov et al, 2000, Anal. Biochem 278:123). In addition, a collection of overexpression clones of yeast proteins have been prepared and screened for biochemical activities (Martzen et al., 1999, Science 286: 1153). However, thousands of individual proteins approximating an entire proteome have not been prepared, arrayed, and screened for multiple activities (Caveman, 2000, J. Cell Sci.113:3543)

Screening an entire proteome would entail the systematic probing of biochemical activities of proteins that are produced in a high throughput fashion, and analyzing the functions of hundreds or thousands of proteins samples in parallel (Zhu et al., 2000, Nat. Genet. 26:283; MacBeath et al., 2000, Science 289:1763; Arenkov et al, 2000, Anal. Biochem 278:123). Attempts to screen an entire proteome array have encountered major obstacles, including the inability to generate the necessary expression clones, and to express and purify the expressed proteins in a high-throughput fashion. *In vitro* assays have previously been conducted using random expression libraries or pooling strategies, both of which have shortcomings (Martzen et al., 1999, Science 286:1153; Bussow et al., 2000, Genomics 65:1). Specifically, random expression libraries are tedious to screen, and contain clones that are often not full-length. Another recent approach has been to generate defined arrays and screen the array using a pooling strategy (Martzen et al. 1999, Science 286:1153). The pooling strategy obscures the actual number of proteins screened, however, and the strategy is cumbersome when large numbers of positives are identified.

Another method useful for detecting protein-protein interactions is the two-hybrid approach (Uetz et al., 2000, Nature 403:623; Ito et al., 2000, Proc. Natl. Acad. Sci. U.S.A.97:1143). The types of interactions that can be detected using this approach are limited, however, because the interactions are typically detected in the nucleus.

Therefore, there remains a need in the art for the large-scale analysis of biochemical functions which would require preparing and screening, in a high-throughput manner, a comprehensive set of proteins encoded by a species's genome.

Citation or identification of any reference in Section 2, or in any other section of this application, shall not be considered an admission that such reference is available as prior art to the present invention.

5 3. SUMMARY OF THE INVENTION

The present invention provides proteome chips useful for the global study of the protein interactions of a species in a high-throughput manner. The methods and compositions of the invention are made possible by Applicants' new and unobvious discovery of a means of preparing a comprehensive set of expression constructs containing protein-coding sequences of a genome, producing the protein products in host cells in a high-throughput fashion, and analyzing the functions of a plurality of proteins in a high-throughput manner using microarrays.

The present invention is directed to proteome chips, which are positionally addressable arrays comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins represents a substantial proportion of all proteins expressed in a single species, wherein translation products of one open reading frame are considered a single protein.

An advantage of using arrays, rather than performing one-by-one assays, is the
ability to identify and characterize many protein-probe interactions simultaneously.

Moreover, complex mixtures of probes can be contacted with a proteome chip to, for example, detect interactions in a milieu more representative of that in a cell, and to quickly evaluate many potential binding compounds.

Accordingly, in one embodiment, the present invention provides a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a single species.

In another embodiment, the invention provides a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40% or 50% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein. In a specific embodiment, the plurality of proteins comprises at least 50% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein.

In another embodiment, the present invention provides a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 500,000 or 1,000,000 protein(s) expressed in a single species.

In another embodiment, the invention provides a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins in aggregate comprise proteins encoded by at least 1000 different known genes in a single species.

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In a further embodiment, the proteins are organized on the array according to a classification of proteins. The classification can be by abundance, function, functional class, enzymatic activity, homology, protein family, association with a particular metabolic or signal transduction pathway, association with a related metabolic or signal transduction pathway, or posttranslational modification.

In another embodiment, the invention provides a positionally addressable array as described above, wherein the solid support comprises glass, ceramics, nitrocellulose, amorphous silicon carbide, castable oxides, polyimides, polymethylmethacrylates, polystyrenes, or silicone elastomers.

In a further embodiment, the solid support comprises a material that helps bind the
plurality of proteins to the solid support. For example, the solid support can be coated with
a material that binds to an affinity tag of each protein. In a particular embodiment, the solid
support comprises glutathione. In another particular embodiment, the solid support coating
comprises nickel or nitrocellulose. In another particular embodiment, the solid support
coating comprises glutathione and nickel. In a one embodiment, the solid support is a
nickel-coated glass slide. In a preferred embodiment, the solid support is a nitrocellulosecoated glass slide. Nitrocellulose-coated glass slides for making protein (and DNA)
microarrays are commercially available (e.g., from Schleicher & Schuell (Keene, NH),
which sells glass slides coated with a nitrocellulose based polymer (Cat. no. 10 484 182)).
In a specific embodiment, each protein is spotted onto the nitrocellulose-coated glass slide
using an OMNIGRIDTM (GeneMachines, San Carlos, CA).

Proteins on the proteome chips preferably are fusion proteins comprising at least one affinity tag useful for purifying and/or attaching the proteins to the proteome chip.

The present invention also provides methods for making proteome chips.

Accordingly, the invention provides a method for constructing a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support,

with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a single species.

In one embodiment, the invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, or 50% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein.

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In another embodiment, the present invention provides a method for constructing a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 500,000 or 1,000,000 protein(s) expressed in a single species.

In another embodiment, the present invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins in aggregate comprise proteins encoded by at least 1000 different known genes in a single species.

The present invention also provides methods for making and isolating viral, prokaryotic or eukaryotic proteins in a readily scalable format, amenable to high-throughput analysis. Preferred methods include synthesizing and purifying proteins in an array format compatible with automation technologies. Accordingly, in one embodiment, the invention provides a method for making and isolating viral, prokaryotic or eukaryotic proteins comprising the steps of growing a eukaryotic cell transformed with a vector having a heterologous sequence operatively linked to a regulatory sequence, contacting the regulatory sequence with an inducer that enhances expression of a protein encoded by the heterologous sequence, lysing the cell, contacting the protein with a binding agent such that a complex between the protein and binding agent is formed, isolating the complex from cellular debris, and isolating the protein from the complex, wherein each step is conducted in a 96-well format.

The protein is preferably a fusion protein such that the heterologous sequence comprises the coding region for the protein of interest and sequences encoding a tag, such as an affinity tag. Such tags can be useful for monitoring the protein, separating the fusion

protein from cellular debris and contaminating reagents, and/or attaching the protein to a proteome chip of the invention.

The present invention also provides methods for making a positionally addressable arrays comprising the step of attaching a plurality of fusion proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the protein comprises a first tag, a second tag, and a protein encoded by a genomic nucleic acid of an organism. In certain embodiments, the protein is tagged with one tag at the aminoterminal end of the protein, and with a second, different tag at the carboxy-terminal end. In other embodiments, the protein is tagged with two tags at the amino-terminal end of the 10 protein, or with two tags at the carboxy-terminal end. In yet other embodiments, the protein is tagged with one or more tags at site(s) on the protein other than the amino- or carboxyterminal end. The advantages of using double-tagged proteins include the ability to obtain highly purified proteins, as well as providing a streamlined manner of purifying proteins from cellular debris and attaching the proteins to a solid support.

Accordingly, in a particular embodiment, the first tag is a glutathione-S-transferase tag ("GST tag") and the second tag is a poly-histidine tag ("His tag"). In another embodiment, the GST tag and the His tag are attached to the amino-terminal end of the protein. Alternatively, the GST tag and the His tag are attached to the carboxy-terminal end of the protein. The GST tag and His tag can be found on either the amino-terminal or 20 carboxy-terminal end of the protein. In certain embodiments, the GST tag is attached to the amino-terminal end of the protein and the His tag is attached to the carboxy-terminal end. In other embodiments, the His tag is attached to the amino-terminal end of the protein and the GST tag is attached to the carboxy-terminal end.

Alternating the placement of an affinity tag of a fusion protein can lead to functional secondary structure, proper folding of extracellular domains, and appropriate trafficking, localization, and/or secretion of proteins. For example, fusion of a GST tag and a His tag onto the carboxy-terminal end of the protein can obviate inappropriate folding or expression when the regions upstream of the translational initiation codon are blocked.

The present invention also provides methods of using a protein array to screen for 30 lipid-binding proteins. Accordingly, in one embodiment, the invention is a method for using a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, comprising the steps of contacting a probe with the array, and detecting protein-probe interaction, wherein the probe comprises a lipid. In a particular embodiment, the lipid comprises a phospholipid such as, but not limited to,

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phosphatidylcholine and phosphatidylinositol. In another particular embodiment, the probe is in the form of a liposome containing phospholipids of interest.

Also provided are methods for using proteome chips. The proteome chips of the invention can be used to assay for essentially all protein-protein interactions in a species or cell. The proteome chips can also be used to systematically assay all the proteins of a species that interact with a test compound. Therefore, using the proteome chips of the invention, a multitude of activities can be assayed to yield a wealth of information such as, but not limited to, defining a "fingerprint" or "signature" of a cell or organism in response to a stimulus, characterizing all proteins in a species that interact with a probe of interest, 10 characterizing all proteins in two or more species that interact with a probe of interest, characterizing all proteins involved in a biological pathway (e.g., metabolic or signal transduction pathway) or in related biological pathways, characterizing all proteins in a species with enzymatic activity(ies) of interest (e.g., kinase activity, protease activity, phosphatase activity, glycosidase, acetylase activity, and other chemical group transferring 15 enzymatic activity), characterizing all proteins in a species with posttranslational modification(s) of interest, and identifying drug targets. In a specific embodiment, proteome chips of the invention are used to characterize all proteins, e.g., drug targets, in a species that interact to with a drug or drug candidate of interest.

Thus, the invention encompasses a method for detecting a binding protein comprising the steps of contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least one protein encoded by at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of the known genes in a single species, and detecting any protein-probe interaction.

In another embodiment, the invention encompasses a method for detecting a binding protein comprising the steps of contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein, and detecting any protein-probe interaction.

In another embodiment, the invention encompasses a method for detecting a binding protein comprising the steps of contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50,

100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 500,000 or 1,000,000 known proteins expressed in a single species, and detecting any protein-probe interaction.

In another embodiment, the invention encompasses a method for detecting a binding protein comprising the steps of contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins in aggregate comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 5000, 10000, 20000, 30000, 40000, or 50000 different known genes in a single species, and detecting any protein-probe interaction.

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In another embodiment, the invention encompasses a method for detecting a binding protein comprising the steps of contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins in aggregate comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 5000, 10000, 20000, 30000, 40000, or 50000 different known genes in at least two species, and detecting any protein-probe interaction.

In another embodiment, the invention encompasses a method for detecting a binding protein comprising the steps of contacting a probe with a positionally addressable array comprising a plurality of fusion proteins, with each protein being at a different position on a solid support, wherein the fusion protein comprises a first tag, a second tag, and a protein sequence encoded by genomic nucleic acid of an organism, and detecting any protein-probe interaction. As described above, in certain embodiments, the two tags can be His and GST.

The present invention also provides a method of labeling a protein for use in a binding assay, comprising the steps of contacting separate aliquots of the protein with a biotin-transferring compound under conditions and for a period of time to produce proteins that are biotinylated to differing degrees among the different aliquots, and combining together the different aliquots to produce a sample of differentially biotinylated protein.

The present invention also provides a method for detecting a binding protein comprising the steps of contacting a sample of biotinylated protein produced by the method described above with a positionally addressable array comprising a plurality of proteins,

with each protein being at a different position on a solid support, and detecting any positions on the array protein-probe interaction wherein interaction between a biotinylated protein and a protein on the array occurs.

The present invention also provides a method for detecting a binding protein comprising the steps of contacting a sample of biotinylated proteins produced by the method described above with a positionally addressable array comprising a plurality of proteins,

with each protein being at a different position on a solid support, contacting the array with streptavidin conjugated to a fluor and detecting positions on the array at which the fluorescence occurs, wherein the fluorescence indicates that interaction between a biotinylated protein and a protein on the array occurs.

The present invention also provides a method for identifying whether a signal is positive. Accordingly, one embodiment of the invention provides a method for identifying whether a signal obtained in an assay using a protein microarray is positive, indicating binding of a probe to an interactor.

10 3.1 Definitions

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As used herein, the word "protein" refers to a full-length protein, a portion of a protein, or a peptide. Proteins can be produced via fragmentation of larger proteins, or chemically synthesized. Preferably, proteins are prepared by recombinant overexpression in a species such as, but not limited to, bacteria, yeast, insect cells, and mammalian cells. Proteins to be placed in a protein microarray of the invention preferably are fusion proteins, more preferably with at least one affinity tag to aid in purification and/or immobilization.

As used herein, the word "interactor" refers to a protein on a protein microarray that interacts with a probe.

As used herein, the word "probe" refers to any chemical reagent such as, but not limited to, a protein, nucleic acid (*e.g.*, DNA, RNA, oligonucleotide, polynucleotide), small molecule, substrate, inhibitor, drug or drug candidate, receptor, antigen, hormone, steroid, lipid, phospholipid, liposome, antibody, cofactor, cytokine, glutathione, immunoglobulin domain, carbohydrate, maltose, nickel, dihydrotrypsin, calmodulin, biotin, lectin, and heavy metal, that can be applied to a protein microarray of the invention to assay for interaction with a protein of the microarray.

4. BRIEF DESCRIPTION OF THE FIGURES

FIGS. 1A-1B. The procedure of yeast proteome analysis using protein chip technology.

A. The yeast ORFs were cloned into a double-tagged yeast *GAL1* expression vector via a recombination strategy and verified for correct identities by sequencing. Each pure plasmid construct was then reintroduced into a yeast strain for large-scale protein purification. Yeast cultures were grown in a 96-well format, and induced by adding

galactose. After the high-throughput purification step, the purified proteins were aliquoted and stored in a glycerol buffer at -80°C before printing. Using a high precision microarrayer, 6566 protein samples can be spotted in duplicate onto 80 slides in a single experiment.

5 B. Immunoblot analysis of proteins purified from 3-ml yeast cultures.

FIGS. 2A-2C. A proteome chip probed with anti-GST antibodies.

Sixty samples were examined by immunoblot analysis using anti-GST antibodies; 19 representative examples are shown. Greater than 80% of the preparations produce high vields of fusion protein.

- A. GST::yeast fusion proteins were purified in a 96-well format. 6566 protein samples representing 5800 unique proteins were spotted in duplicate on a single nickel-coated microscope slide. The slide was probed with anti-GST antibodies.
- B. An enlarged image of one of the 48 blocks is depicted to the right of the proteome chip. The letters indicate the duplicated protein samples, and the numbers represent the source plate numbers. Note the excellent resolution and high signal-to-noise ratios.
 - C. Distribution of errors between duplicated spots of each ORF product. The unit of ϵ is the signal intensity determined by the Axon scanner. As determined by a GST control, ten units are equivalent to approximately 32 fg of protein. The center of the ϵ distribution is 1.2 units, and 95% of the samples are within 10 units from the center.

FIGS. 3A-3B. Examples of different assays on the proteome chips.

A. Proteome chips containing 6566 yeast proteins were spotted in duplicate and incubated with the biotinylated probes indicated. Positive interactors, indicated by boxes, were identified in six protein-liposome interaction assays ("PI(3)P", "PI(4,5)P₂", "PI(4)P", "PI(3,4)P₂", "PI(3,4,5)P₃", "PC"), a calmodulin-binding assay ("Calmodulin"), and a DNA-protein interaction assay ("Genomic DNA"). Each block contains 16X18 protein spots. The positive signals in duplicate appear as horizontal pairs in the bottom panels. The duplicate spotting serves as an internal control, which is important when the signals are weak relative to the background. The upper panels show the same yeast protein preparations of a control proteome chip probed with anti-GST antibodies ("α-GST"). As demonstrated by the figure, strong signals are often observed in samples having relatively low levels of GST fusion protein ("Probe"), indicating that the binding is sensitive and specific. PI, phosphatidylinositol. PC, phosphatidylcholine.

B. A putative calmodulin-binding motif identified by searching for amino acid sequences that are shared by the different calmodulin targets (Zhu et al. 2000, Nat. Genet. 26:283). 14 of 39 positive proteins share a motif whose consensus is I/L-Q-X-K-K/X-G-B (SEO ID NO: 1), where X is any residue and B is a basic residue. The size of the lettering, depicted above the alignment, indicates the relative frequency of the amino acid indicated. YFL003C/MSH4 (SEQ ID NO: 2); YJR073C/OPI3 (SEQ ID NO: 3); YBR050C/REG2 (SEO ID NO: 4); YNL202W/SPS19 (SEQ ID NO: 5); YOL016C/CMK2 (SEQ ID NO: 6); BR011C/IPP1 (SEO ID NO: 7); YGR034W/RPL26B (SEQ ID NO: 8); YFR004W/RPN11 (SEO ID NO: 9); YIL021W/RPB3 (SEQ ID NO: 10); YGL063W/PUS2 (SEQ ID NO: 11); 10 YDR292C/SRP101 (SEQ ID NO: 12); YFR014C/CMK1 (SEQ ID NO: 13); YBR213W/MET8 (SEQ ID NO: 14); YAL029C/MYO4 (SEQ ID NO: 15).

FIGS. 4A-4D. Analysis of the phosphatidylinositol-binding proteins. To determine the phosphatidylinositol ("PI")-binding specificity of 150 positive proteins, their binding 15 signals were normalized against the corresponding binding signals of phosphatidylcholine ("PC"). Based on the ratios ("PI/PC"), the proteins were grouped into four categories: (A) 30 strong and specific, (B) 43 strong and nonspecific, (C) 19 weak and specific, and (D) 58 weak and nonspecific phosphatidylinositol-binding proteins. The intensity represents the PI/PC signal ratio as shown by the scale in the figure. The column, labeled "10" to the 20 right of the PI/PC binding ratios indicates the maximum binding signal intensity (open boxes) and its confidence interval (solid horizontal lines); the numbers indicate the log of the values. Boxes in the three columns to the right of the confidence interval column indicate membrane-associated proteins ("Membrane"), kinases ("Kinase"), and uncharacterized ORFs ("Unknown"), respectively.

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- FIGS. 5A-5C. Conventional methods confirm protein-lipid interactions detected by the proteome microarrays (Casamayor et al., 1999, Curr. Biol. 9:186; Guerra et al., 2000, Biosci. Rep. 20:41).
- A. PI(4.5)P2 liposomes were first adhered to a nitrocellulose membrane, which was 30 blocked by BSA; a dilution series of Rim15p, Eno2p, and Hxk1p, and a GST control were used to probe the membrane. The bound proteins were detected using the anti-GST antibodies and an enhanced chemiluminescence ("ECL") kit.
- B. A reverse assay was carried out to test for protein-lipid interactions. The proteins were prepared and spotted onto nitrocellulose filters in a dilution series and probed 35 with the six different liposomes. As a control, the six liposomes were also added to the

BSA-blocked membrane. After extensive washing, the bound liposomes were detected using an HRP-conjugated streptavidin and an ECL kit.

C. Linear correlation between the binding signals and the amounts of Rim15p in a membrane assay. When liposome-binding signals of Rim15p from the membrane assay (FIG. 4B) were plotted against the concentration gradient of the spotted Rim15p, the binding signals of PI(4)P, PI(3,4)P2, PI(3)P, and PI(4,5)P2 correlated linearly with the amounts of Rim15p. The interaction of PI(4)P and Rim15p showed the highest affinity, which was at least three-fold higher than the affinity of the control PC with Rim15p.

10 5. DETAILED DESCRIPTION OF THE INVENTION

The present invention is based, in part, on Applicants' construction of a yeast proteome microarray containing approximately 80% of all proteins expressed in yeast, representing the first description of a proteome array for any species. The proteome chips of the invention can be used for global analyses of protein interactions and activities in a species. The use of proteome chips has significant advantages over existing approaches. One advantage of the proteome microarray technology presented here is that a large set of individual proteins can be directly screened in a high-throughput fashion for hundreds or even thousands of biochemical activities simultaneously. For example, an advantage of the proteome chip approach is that proteins can be directly screened *in vitro* for a wide variety of activities including, but not limited to, protein-drug interactions (*e.g.*, drug target-drug interactions), protein-lipid interactions and enzymatic assays. In addition, a wide range of *in vitro* conditions can be readily tested. Furthermore, once the proteins are prepared, proteome screening is significantly faster and cheaper, and rapid data analysis in many microarray formats is compatible with existing equipment and analytical software.

5.1 Proteome Arrays.

The present invention encompasses a positionally addressable arrays comprising a plurality of proteins, with each protein being at a different position on a solid support,

wherein the plurality of proteins comprises at least one protein encoded by at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of the known genes in a single species, *i.e.*, all protein isoforms and splice variants derived from a gene are considered one protein.

The present invention encompasses a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support,

wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a single species, where protein isoforms and splice variants are counted as a single protein. In one embodiment, the plurality of proteins comprises about 90%, 95%, or 99% of all proteins expressed in a species. In a particular embodiment, the plurality of proteins comprises about 93.5% of all proteins expressed in a species.

The present invention also encompasses a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 500,000 or 1,000,000 protein(s) expressed in a single species.

The present invention also encompasses a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins in aggregate comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 5000, 10000, 20000, 30000, 40000, or 50000 different known genes in a single species.

The present invention also encompasses a positionally addressable array comprising a plurality of fusion proteins to a surface of a solid support, with each fusion protein being at a different position on the solid support, wherein the fusion protein comprises a first tag, a second tag, and a protein sequence encoded by genomic nucleic acid of an organism. In another embodiment, the protein sequence of the fusion protein need not be encoded in a genomic nucleic acid of an organism, but is a sequence for which it is desired to identify a function and/or activity of a binding protein.

A positionally addressable array provides a configuration such that each probe or protein of interest is at a known position on the solid support thereby allowing the identity of each probe or protein to be determined from its position on the array. Accordingly, each protein on an array is preferably located at a known, predetermined position on the solid support such that the identity of each protein can be determined from its position on the solid support.

In one embodiment, the species is a virus. In another embodiment, the species is a prokaryote. In another embodiment, the species is a eukaryote. In another embodiment, the species is a vertebrate. In yet another embodiment, the species is a mammal. In a particular embodiment, the species is an animal, including, but not limited to, an insect, primate, and rodent. In a specific embodiment, the species is a monkey, fruit fly, cow, horse, sheep, pig, chicken, turkey, quail, cat, dog, mouse, rat, rabbit, nematode or fish. In a preferred

embodiment, the species is a human. In another preferred embodiment, the species is a yeast.

Proteins of the proteome chips of the invention include full-length proteins, portions of full-length proteins, and peptides, which can be prepared by recombinant overexpression, fragmentation of larger proteins, or chemical synthesis. Proteins can be overexpressed in cells derived from, for example, yeast, bacteria, insects, humans, or non-human mammals such as mice, rats, cats, dogs, pigs, cows and horses. Further, fusion proteins comprising a defined domain attached to a natural or synthetic protein can be used. Proteins of the proteome chips can be purified prior to being attached to the solid support of the chip. Also 10 the proteins of the proteome chips can be purified, or further purified, during attachment to the proteome chip.

Proteins can be embedded in artificial or natural membranes (e.g., liposomes, membrane vesicles) prior to, or at the time of attachment to the protein chip. In fact, the synthesis of certain proteins may preferably be conducted in the presence of artificial or 15 natural membranes to, for example, promote protein folding, protein processing, retain activity, and/or prevent precipitation of the protein.

Further, proteins can be attached to the solid support of the proteome chip. Alternatively, the proteins can be delivered into wells of the proteome chip, where they remain unbound to the solid support of the proteome chip.

20 The present invention is also directed to compounds useful as solid supports for the proteome chips of the invention. The solid support can be constructed from materials such as, but not limited to, silicon, glass, quartz, polyimide, acrylic, polymethylmethacrylate (LUCITE®), ceramic, nitrocellulose, amorphous silicon carbide, polystyrene, and/or any other material suitable for microfabrication, microlithography, or casting. For example, the 25 solid support can be a hydrophilic microtiter plate (e.g., MILLIPORETM) or a nitrocellulosecoated glass slide. In a preferred embodiment, the solid support is a nitrocellulose-coated glass slide. Nitrocellulose-coated glass slides for making protein (and DNA) microarrays are commercially available (e.g., from Schleicher & Schuell (Keene, NH), which sells glass slides coated with a nitrocellulose based polymer (Cat. no. 10 484 182)). In a specific 30 embodiment, each protein is spotted onto the nitrocellulose-coated glass slide using an OMNIGRIDTM (GeneMachines, San Carlos, CA). The present invention contemplates other solid supports useful for constructing a protein chip, some of which are disclosed, for example, in co-pending United States Application No. 09/849,781, which was filed on May 4, 2001, and which is incorporated herein by reference in its entirety.

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In a particular embodiment, the solid support comprises a silicone elastomeric material such as, but not limited to, polydimethylsiloxane ("PDMS"). An advantage of silicone elastomeric materials is their flexible nature.

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In another particular embodiment, the solid support is a silicon wafer. The silicon wafer can be patterned and etched (*see*, *e.g.*, G. Kovacs, 1998, <u>Micromachined Transducers Sourcebook</u>, Academic Press; M. Madou, 1997, <u>Fundamentals of Microfabrication</u>, CRC Press. The etched wafer can be used to cast the proteome chips of the invention.

In one embodiment, the present invention provides a proteome chip comprising a solid support that is a flat surface such as, but not limited to, a glass slide. Dense protein arrays can be produced on, for example, glass slides, such that assays for the presence, amount, and/or functionality of proteins can be conducted in a high-throughput manner.

Accordingly, in one embodiment, the proteome chip comprises a plurality of proteins that are applied to the surface of a solid support, wherein the density of the sites at which protein are applied is at least 100 sites/cm², 1000 sites/cm², 10,000 sites/cm², 1000,000 sites/cm², 10,000,000 sites/cm², 10,000,000 sites/cm², 25,000,000 sites/cm², 10,000,000,000 sites/cm². Each individual protein sample is preferably applied to a separate site on the chip. The identity of the protein(s) at each site on the chip is/are known.

In another embodiment, the solid support has an array of wells. The use of microlithographic and micromachining fabrication techniques (*see*, *e.g.*, co-pending United States Application No. 09/849,781, filed on May 4, 2001, which is incorporated herein by reference in its entirety) can be used to create well arrays with a wide variety of dimensions ranging from hundreds of microns down to 100 nm or even smaller, with well depths of similar dimensions. In one embodiment, a silicon wafer is micromachined and acts as a master mold to cast wells of 400 μm diameter that are spaced 200 μm apart, for a well density of about 277 wells per cm², with individual well volumes of about 30 nl for 100 μm deep wells.

In another embodiment, microlithographic micromachining is used to fabricate wells 500 nm and 275 nm diameter, spaced 1 µm apart to yield well densities of over 44 million and over 61 million wells per cm² respectively. Higher densities are possible through closer spacing, as well as through smaller diameters.

In another embodiment, precision laser micromachining techniques can be used to directly fabricate mold structures out of acrylic with dimensions ranging from greater than 1.5 mm down to 500 μ m, with well spacing of about 500 μ m. Volumes of these wells are in the 50-500 nl range.

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Accordingly, in one embodiment, the proteome chip comprises a plurality of wells on the surface of a solid support, wherein the density of wells is at least 100 wells/cm², 1000 wells/cm², 10,000 wells/cm², 100,000 wells/cm², 1,000,000 wells/cm², 10,000,000 wells/cm², 25,000,000 wells/cm², 10,000,000,000 wells/cm², or 10,000,000,000,000 wells/cm². The present invention contemplates variations of protein chips comprising a plurality of wells, which are disclosed, for example, in co-pending United States Application No. 09/849,781, filed on May 4, 2001, and which is incorporated herein by reference in its entirety.

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The present invention also contemplates variations in the shape, width-to-depth 10 ratio and volume of wells in the proteome chip, which are disclosed, for example, in co-pending United States Application No. 09/849,781, filed on May 4, 2001, and which is incorporated herein by reference in its entirety. Such shapes include, but are not limited to circular, oval, rectangular, square, etc. The wells can also have, for example, square, round V-shaped or U-shaped bottoms.

In one embodiment, the solid support comprises gold. In a preferred embodiment, the solid support comprises a gold-coated slide. In another embodiment, the solid support comprises nickel. In another preferred embodiment, the solid support comprises a nickel-coated slide. Solid supports comprising nickel are advantageous for purifying and attaching fusion proteins having a poly-histidine tag ("His tag"). In another embodiment, 20 the solid support comprises nitrocellulose. In another preferred embodiment, the solid support comprises a nitrocellulose-coated slide.

The invention further relates to compounds useful for derivatization of the proteome chip substrate. The proteins can be bound directly to the solid support, or can be attached to the solid support through a linker molecule or compound. The linker can be any molecule 25 or compound that derivatizes the surface of the solid support to facilitate the attachment of proteins to the surface of the solid support. The linker may covalently or non-covalently bind the proteins or probes to the surface of the solid support. In addition, the linker can be an inorganic or organic molecule. In certain embodiments, the linker may be a silane, e.g., sianosilane, thiosilane, aminosilane, etc. The present invention contemplates compounds 30 useful for derivatization of a protein chip, some of which are disclosed, for example, in co-pending United States Application No. 09/849,781, which was filed on May 4, 2001, and which is incorporated herein by reference in its entirety.

Accordingly, in one embodiment, the proteins of the proteome chip are bound non-covalently to the solid support (e.g., by adsorption). Proteins that are non-covalently bound to the solid support can be attached to the surface of the solid support by a variety of

molecular interactions such as, for example, hydrogen bonding, van der Waals bonding, electrostatic, or metal-chelate coordinate bonding. In a particular embodiment, proteins are bound to a poly-lysine coated surface of the solid support. In addition, as described above, in certain embodiments, the proteins are bound to a silane (*e.g.*, sianosilane, thiosilane, aminosilane, etc.) coated surface of the solid support.

In addition, crosslinking compounds commonly known in the art, *e.g.* homo- or heterofunctional crosslinking compounds (*e.g.*, bis[sulfosuccinimidyl]suberate, N-[gamma-maleimidobutyryloxy]succinimide ester, or 1-ethyl-3-[3-dimethylaminopropyl]carbodiimide), may be used to attach proteins to the solid support via covalent or non-covalent interactions.

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In another embodiment, the proteins of the proteome chip are bound covalently to the solid support. For example, the proteins can be bound to the solid support by receptor-ligand interactions, which include interactions between antibodies and antigens, DNA-binding proteins and DNA, enzyme and substrate, avidin (or streptavidin) and biotin (or biotinylated molecules), and interactions between lipid-binding proteins and phospholipids (or membranes, vesicles, or liposomes comprising phospholipids).

Purified proteins can be placed on an array using a variety of methods known in the art. In one embodiment, the proteins are printed onto the solid support. In a further embodiment, the proteins are attached to the solid support using an affinity tag. Use of an affinity tag different from that used to purify the proteins is preferred, since further purification is achieved when building the protein array.

Accordingly, in a preferred embodiment, proteins of the proteome chip are expressed as fusion proteins having at least one heterologous domain with an affinity for a compound that is attached to the surface of the solid support. Suitable compounds useful for binding fusion proteins onto the solid support (*i.e.*, acting as binding partners) include, but are not limited to, trypsin/anhydrotrypsin, glutathione, immunoglobulin domains, maltose, nickel, or biotin and its derivatives, which bind to bovine pancreatic trypsin inhibitor, glutathione-S-transferase, Protein A or antigen, maltose binding protein, poly-histidine (*e.g.*, HisX6 tag), and avidin/streptavidin, respectively. For example, Protein A, Protein G and Protein A/G are proteins capable of binding to the Fc portion of mammalian immunoglobulin molecules, especially IgG. These proteins can be covalently coupled to, for example, a Sepharose® support to provide an efficient method of purifying fusion proteins having a tag comprising an Fc domain.

In a further embodiment, the proteins are bound directly to the solid support. In another further embodiment, the proteins are bound to the solid support via a linker. In a

particular embodiment, the proteins are attached to the solid support via a His tag. In another particular embodiment, the proteins are attached to the solid support via a 3-glycidooxypropyltrimethoxysilane ("GPTS") linker. In a specific embodiment, the proteins are bound to the solid support via His tags, wherein the solid support comprises a flat surface. In a preferred embodiment, the proteins are bound to the solid support via His tags, wherein the solid support comprises a nickel-coated glass slide.

The proteome chips of the present invention are not limited in their physical dimensions and can have any dimensions that are useful. Preferably, the proteome chip has an array format compatible with automation technologies, thereby allowing for rapid data analysis. Thus, in one embodiment, the proteome microarray format is compatible with laboratory equipment and/or analytical software. In a preferred embodiment, the proteome chip is the size of a standard microscope slide. In another preferred embodiment, the protein chip is designed to fit into a sample chamber of a mass spectrometer.

15 5.2 Methods for Making and Purifying Proteins in a High Density Array Format.

The present invention also relates to methods for making and isolating viral, prokaryotic or eukaryotic proteins in a readily scalable format, amenable to high-throughput analysis. Preferred methods include synthesizing and purifying proteins in an array format compatible with automation technologies. Accordingly, in one embodiment, the invention provides a method for making and isolating eukaryotic proteins comprising the steps of growing a eukaryotic cell transformed with a vector having a heterologous sequence operatively linked to a regulatory sequence, contacting the regulatory sequence with an inducer that enhances expression of a protein encoded by the heterologous sequence, lysing the cell, contacting the protein with a binding agent such that a complex between the protein and binding agent is formed, isolating the complex from cellular debris, and isolating the protein from the complex, wherein each step is conducted in a 96-well format.

In one embodiment, the invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least one protein encoded by at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of the known genes in a single species.

In another embodiment, the invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a

solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein.

In another embodiment, the invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 10 500,000 or 1,000,000 protein(s) expressed in a single species.

In yet another embodiment, the invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins in aggregate comprises at least 1, 2, 3, 4, 5, 10, 20, 15 30, 40, 50, 100, 200, 500, 1000, 5000, 10000, 20000, 30000, 40000, or 50000 different known genes in a single species.

In yet another embodiment, the invention provides a method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid 20 support, wherein the protein is a fusion protein comprising a first tag, a second tag, and a protein encoded by genomic nucleic acid of an organism.

In one embodiment, each step in the synthesis and purification procedures is conducted in an array amenable to rapid automation. Such arrays can comprise a plurality of wells on the surface of a solid support wherein the density of wells is at least 10, 20, 30, 25 40, 50, 100, 1000, 10,000, 100,000, or 1,000,000 wells/cm², for example. Alternatively, such arrays comprise a plurality of sites on the surface of a solid support, wherein the density of sites is at least 10, 20, 30, 40, 50, 100, 1000, 10,000, 100,000, or 1,000,000 sites/cm², for example.

In a particular embodiment, eukaryotic proteins are made and purified in a 96-array 30 format (i.e., each site on the solid support where processing occurs is one of 96 sites), e.g., in a 96-well microtiter plate. In a preferred embodiment, the solid support does not bind proteins (e.g., a non-protein-binding microtiter plate).

In certain embodiments, proteins are synthesized by in vitro translation according to methods commonly known in the art.

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Any expression construct having an inducible promoter to drive protein synthesis can be used in accordance with the methods of the invention. Preferably, the expression construct is tailored to the cell type to be used for transformation. Compatibility between expression constructs and host cells are known in the art, and use of variants thereof are also encompassed by the invention.

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Any host cell that can be grown in culture can be used to synthesize the proteins of interest. Preferably, host cells are used that can overproduce a protein of interest, resulting in proper synthesis, folding, and posttranslational modification of the protein. Preferably, such protein processing forms epitopes, active sites, binding sites, etc. useful for assays to characterize molecular interactions *in vitro* that are representative of those *in vivo*.

Accordingly, a eukaryotic cell (e.g., yeast, human cells) is preferably used to synthesize eukaryotic proteins. Further, a eukaryotic cell amenable to stable transformation, and having selectable markers for identification and isolation of cells containing transformants of interest, is preferred. Alternatively, a eukaryotic host cell deficient in a gene product is transformed with an expression construct complementing the deficiency. Cells useful for expression of engineered viral, prokaryotic or eukaryotic proteins are known in the art, and variants of such cells can be appreciated by one of ordinary skill in the art.

For example, the InsectSelect system from Invitrogen (Carlsbad, CA, catalog no. K800-01), a non-lytic, single-vector insect expression system that simplifies expression of high-quality proteins and eliminates the need to generate and amplify virus stocks, can be used. A preferred vector in this system is pIB/V5-His TOPO TA vector (catalog no. K890-20). Polymerase chain reaction ("PCR") products can be cloned directly into this vector, using the protocols described by the manufacturer, and the proteins can be expressed with N-terminal histidine tags useful for purifying the expressed protein.

Another eukaryotic expression system in insect cells, the BAC-TO-BACTM system (LIFETECHTM, Rockville, MD), can also be used. Rather than using homologous recombination, the BAC-TO-BACTM system generates recombinant baculovirus by relying on site-specific transposition in *E. coli*. Gene expression is driven by the highly active polyhedrin promoter, and therefore can represent up to 25% of the cellular protein in infected insect cells.

In a particular embodiment, yeast cultures are used to synthesize eukaryotic fusion proteins. Fresh cultures are preferably used for efficient induction of protein synthesis, especially when conducted in small volumes of media. Also, care is preferably taken to prevent overgrowth of the yeast cultures. In addition, yeast cultures of about 3 ml or less are preferable to yield sufficient protein for purification. To improve aeration of the

cultures, the total volume can be divided into several smaller volumes (e.g., four 0.75 ml cultures can be prepared to produce a total volume of 3 ml).

Cells are then contacted with an inducer (e.g., galactose), and harvested. Induced cells are washed with cold (i.e., 4°C to about 15°C) water to stop further growth of the cells, and then washed with cold (i.e., 4°C to about 15°C) lysis buffer to remove the culture medium and to precondition the induced cells for protein purification, respectively. Before protein purification, the induced cells can be stored frozen to protect the proteins from degradation. In a specific embodiment, the induced cells are stored in a semi-dried state at -80°C to prevent or inhibit protein degradation.

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Cells can be transferred from one array to another using any suitable mechanical device. For example, arrays containing growth media can be inoculated with the cells of interest using an automatic handling system (e.g., automatic pipette). In a particular embodiment, 96-well arrays containing a growth medium comprising agar can be inoculated with yeast cells using a 96-pronger. Similarly, transfer of liquids (e.g., reagents) from one array to another can be accomplished using an automated liquid-handling device (e.g., Q-FILLTM, Genetix, UK).

Although proteins can be harvested from cells at any point in the cell cycle, cells are preferably isolated during logarithmic phase when protein synthesis is enhanced. For example, yeast cells can be harvested between OD₆₀₀=0.3 and OD₆₀₀=1.5, preferably between OD₆₀₀=0.5 and OD₆₀₀=1.5. In a particular embodiment, proteins are harvested from the cells at a point after mid-log phase. Harvested cells can be stored frozen for future manipulation.

The harvested cells can be lysed by a variety of methods known in the art, including mechanical force, enzymatic digestion, and chemical treatment. The method of lysis should be suited to the type of host cell. For example, a lysis buffer containing fresh protease inhibitors is added to yeast cells, along with an agent that disrupts the cell wall (e.g., sand, glass beads, zirconia beads), after which the mixture is shaken violently using a shaker (e.g., vortexer, paint shaker).

In a specific embodiment, zirconia beads are contacted with the yeast cells, and the cells lysed by mechanical disruption by vortexing. In a further embodiment, lysing of the yeast cells in a high-density array format is accomplished using a paint shaker. The paint shaker has a platform that can firmly hold at least eighteen 96-well boxes in three layers, thereby allowing for high-throughput processing of the cultures. Further the paint shaker violently agitates the cultures, even before they are completely thawed, resulting in efficient disruption of the cells while minimizing protein degradation. In fact, as determined by

microscopic observation, greater than 90% of the yeast cells can be lysed in under two minutes of shaking.

The resulting cellular debris can be separated from the protein and/or other molecules of interest by centrifugation. Additionally, to increase purity of the protein sample in a high-throughput fashion, the protein-enriched supernatant can be filtered, preferably using a filter on a non-protein-binding solid support. To separate the soluble fraction, which contains the proteins of interest, from the insoluble fraction, use of a filter plate is highly preferred to reduce or avoid protein degradation. Further, these steps preferably are repeated on the fraction containing the cellular debris to increase the yield of protein.

Proteins can then be purified from the protein-enriched supernatant using a variety of affinity purification methods known in the art. Affinity tags useful for affinity purification of fusion proteins by contacting the fusion protein preparation with the binding partner to the affinity tag, include, but are not limited to, calmodulin,

trypsin/anhydrotrypsin, glutathione, immunoglobulin domains, maltose, nickel, or biotin and its derivatives, which bind to calmodulin-binding protein, bovine pancreatic trypsin inhibitor, glutathione-S-transferase ("GST tag"), antigen or Protein A, maltose binding protein, poly-histidine ("His tag"), and avidin/streptavidin, respectively. Other affinity tags can be, for example, myc or FLAG. Fusion proteins can be affinity purified using an appropriate binding compound (i.e., binding partner such as a glutathione bead), and isolated by, for example, capturing the complex containing bound proteins on a non-protein-binding filter. Placing one affinity tag on one end of the protein (e.g., the carboxy-terminal end), and a second affinity tag on the other end of the protein (e.g., the amino-terminal end) can aid in purifying full-length proteins.

In a particular embodiment, the fusion proteins have GST tags and are affinity purified by contacting the proteins with glutathione beads. In further embodiment, the glutathione beads, with fusion proteins attached, can be washed in a 96-well box without using a filter plate to ease handling of the samples and prevent cross contamination of the samples.

In addition, fusion proteins can be eluted from the binding compound (e.g., glutathione bead) with elution buffer to provide a desired protein concentration. In a specific embodiment, fusion proteins are eluted from the glutathione beads with 30 μ l of elution buffer to provide a desired protein concentration.

For purified proteins that will eventually be spotted onto microscope slides, the glutathione beads are separated from the purified proteins. Preferably, all of the glutathione

beads are removed to avoid blocking of the microarrays pins used to spot the purified proteins onto a solid support. In a preferred embodiment, the glutathione beads are separated from the purified proteins using a filter plate, preferably comprising a non-protein-binding solid support. Filtration of the eluate containing the purified proteins should result in greater than 90% recovery of the proteins.

The elution buffer preferably comprises a liquid of high viscosity such as, for example, 15% to 50% glycerol, preferably about 40% glycerol. The glycerol solution stabilizes the proteins in solution, and prevents dehydration of the protein solution during the printing step using a microarrayer.

Purified proteins are preferably stored in a medium that stabilizes the proteins and prevents dessication of the sample. For example, purified proteins can be stored in a liquid of high viscosity such as, for example, 15% to 50% glycerol, preferably in about 40% glycerol. It is preferred to aliquot samples containing the purified proteins, so as to avoid loss of protein activity caused by freeze/thaw cycles.

The skilled artisan can appreciate that the purification protocol can be adjusted to control the level of protein purity desired. In some instances, isolation of molecules that associate with the protein of interest is desired. For example, dimers, trimers, or higher order homotypic or heterotypic complexes comprising an overproduced protein of interest can be isolated using the purification methods provided herein, or modifications thereof.

Furthermore, associated molecules can be individually isolated and identified using methods

5.3 Methods for Making a Proteome Array.

known in the art (e.g., mass spectroscopy).

The present invention is also directed to methods of making proteome chips.

25 Accordingly, the invention provides methods for constructing a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least one representative protein for at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of the known genes in a species, wherein the protein is all protein isoforms and splice variants derived from a gene.

In another embodiment, the invention provides methods for constructing a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%,

20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a species.

In another embodiment, the present invention provides a method for constructing a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 500,000 or 1,000,000 protein(s) expressed in a species.

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The present invention also relates to methods for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the solid support is cast from a microfabricated mold, and wherein the plurality of proteins comprises at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of all proteins expressed in a species, or comprises at least 1, 2, 3, 4, 5, 10, 20, 30, 40, 50, 100, 200, 500, 1000, 1500, 2000, 2500, 3000, 4000, 5000, 6000, 7000, 8000, 9000, 10,000, 100,000, 500,000 or 1,000,000 protein(s) expressed in a species, or comprises at least one representative protein for at least 1%, 2%, 3%, 4%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 99% of the known genes in a species, wherein the protein is all protein isoforms and splice variants derived from a gene. The present invention contemplates a variety of solid supports cast from a microfabricated mold, some of which are disclosed, for example, in co-pending United States Application No. 09/849,781, filed on May 4, 2001, which is incorporated herein by reference in its entirety.

The present invention also relates to methods for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the protein comprises a first tag and a second tag. The advantages of using double-tagged proteins include the ability to obtain highly purified proteins, as well as providing a streamlined manner of purifying proteins from cellular debris and attaching the proteins to a solid support. In a particular embodiment, the first tag is a glutathione-S-transferase tag

("GST tag") and the second tag is a poly-histidine tag ("His tag"). In a further embodiment, the GST tag and the His tag are attached to the amino-terminal end of the protein.

Alternatively, the GST tag and the His tag are attached to the carboxy-terminal end of the protein.

In yet another embodiment, the GST tag is attached to the amino-terminal end of the protein. In a further embodiment, the His tag is attached to the carboxy-terminal end of the

protein. In yet another embodiment, the His tag is attached to the amino-terminal end of the protein. In a further embodiment, the GST tag is attached to the carboxy-terminal end of the protein.

In yet another embodiment, the protein comprises a GST tag and a His tag, and neither the GST tag nor the His tag is located at the amino-terminal or carboxy-terminal end of the protein. In a specific embodiment, the GST tag and His tag are located within the coding region of the protein of interest; preferably in a region of the protein not affecting the binding domain of interest.

In one embodiment, the first tag is used to purify a fusion protein. In another embodiment, the second tag is used to attach a fusion protein to a solid support. In a specific further embodiment, the first tag is a GST tag and the second tag is a His tag.

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The protein preferably is a fusion protein such that the heterologous sequence comprises the coding region for the protein of interest and sequences encoding a tag, such as an affinity tag. Such tags can be useful for monitoring the protein, separating the fusion protein from cellular debris and contaminating reagents, and/or attaching the protein to a proteome chip of the invention.

Examples of inducers include, but are not limited to, galactose, enhancer-binding proteins, and other transcription factors. In one embodiment, galactose is contacted with a regulatory sequence comprising a galactose-inducible *GAL1* promoter.

A binding agent that can be used in accordance with the invention includes, but is not limited to, a glutathione bead, a nickel-coated solid support, and an antibody. In one embodiment, the complex comprises a fusion protein having a GST tag bound to a glutathione bead. In another embodiment, the complex comprises the a fusion protein having a His tag bound to a nickel-coated solid support. In yet another embodiment, the complex comprises the protein of interest bound to an antibody and, optionally, a secondary antibody.

5.4 Methods for Using a Proteome Array.

The invention is also directed to methods for using proteome chips to assay the presence, amount, and/or functionality of proteins present in at least one sample. Using the proteome chips of the invention, chemical reactions and assays in a large-scale parallel analysis can be performed to characterize biological states or biological responses, and determine the presence, amount, and/or biological activity of proteins. Accordingly, the proteome chips of the invention can be used to assay for essentially all protein-protein interactions in a cell, tissue, organ, system, or organism.

Additionally, the proteome chips of the invention can be used to assay for biological responses to a particular stimulus given to a host cell (i.e., a cell used to produce the fusion protein). For example, yeast cells transformed with expression vectors encoding fusion proteins can be subjected to a stimulus, after which the fusion proteins are purified and arrayed. The arrayed proteins can then be characterized by, for example, probing with any binder. The resulting binding pattern is then compared to an identical array produced from yeast cells not subjected to the stimulus, or subjected to a different stimulus. Differences in the binding patterns can be characteristic of the biological response, and can identify specific interactors of interest with respect to the biological response.

10 In one embodiment, proteome chips prepared from host cells, each chip representing host cells exposed to a different stimulus, are screened with a labeled lectin (e.g., concanavalin A). The pattern of protein-probe interactions, indicating the presence of glycosylated proteins, is compared to determine the effect of each stimulus on the glycosylation state of the proteins of the proteome chip.

Biological activity that can be determined using a proteome chip of the invention includes, but is not limited to, enzymatic activity (e.g., kinase activity, protease activity, phosphatase activity, glycosidase, acetylase activity, and other chemical group transferring enzymatic activity), nucleic acid binding, hormone binding, etc. High density and small volume chemical reactions can be advantageous for the methods relating to using the 20 proteome chips of the invention.

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Further information regarding biological states or responses can be obtained using the proteome chips of the invention, wherein proteins on the chip are organized according to a classification of proteins. The classification can be by abundance, function, functional class, enzymatic activity, homology, protein family, association with a particular metabolic or signal transduction pathway, association with a related metabolic or signal transduction pathway, or posttranslational modification.

Upon contacting the proteins of a proteome chip of the invention with one or more probes, protein-probe interactions can be assayed using a variety of techniques known in the art. For example, the proteome chip can be assayed using standard enzymatic assays that 30 produce chemiluminescence or fluorescence. Various protein modifications can be detected by, for example, photoluminescence, chemiluminescence, or fluorescence using non-protein substrates, enzymatic color development, mass spectroscopic signature markers, or amplification of oligonucleotide tags.

The probe is labeled or tagged with a marker so that its binding can be detected, 35 directly or indirectly, by methods commonly known in the art. Any art-known marker may

be used, including but not limited to tags such as epitope tags, haptens, and affinity tags, antibodies, labels, etc., providing that it is not the same as the affinity tag or reagent used to attach the protein(s) of the proteome chip to the solid substrate of the chip. For example, if biotin is used as a linker to attach proteins to a proteome chip array, then another tag not present in the protein(s) of the proteome chip, *e.g.*, His or GST, is used to label the probe and to detect a protein-probe interaction. In certain embodiments, a photoluminescent, chemiluminescent, fluorescent, or enzymatic tag is used. In other embodiments, a mass spectroscopic signature marker is used. In yet other embodiments, an amplifiable oligonucleotide, peptide or molecular mass label is used.

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In a specific embodiment, the invention provides a method for detecting a proteinprobe interaction comprising the steps of contacting a sample of labeled probe (e.g., labeled protein) with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support; and detecting any positions on the array wherein interaction between the labeled probe and a protein on the array occurs.

15 Accordingly, protein-probe interactions can be detected by, for example, 1) using radioactively labeled ligand followed by autoradiography and/or phosphoimager analysis; 2) binding of hapten, which is then detected by a fluorescently labeled or enzymatically labeled antibody or high-affinity hapten ligand such as biotin or streptavidin; 3) mass spectrometry; 4) atomic force microscopy; 5) fluorescent polarization methods; 6) infrared 20 red labeled compounds or proteins; 7) amplifiable oligonucleotides, peptides or molecular mass labels; 8) stimulation or inhibition of the protein's enzymatic activity; 9) rolling circle amplification-detection methods (Hatch et al., 1999, "Rolling circle amplification of DNA immobilized on solid surfaces and its application to multiplex mutation detection", Genet. Anal. 15:35-40); 10) competitive PCR (Fini et al., 1999, "Development of a 25 chemiluminescence competitive PCR for the detection and quantification of parvovirus B19 DNA using a microplate luminometer", Clin Chem. 45:1391-6; Kruse et al., 1999, "Detection and quantitative measurement of transforming growth factor-beta1 (TGF-beta1) gene expression using a semi-nested competitive PCR assay", Cytokine 11:179-85; Guenthner and Hart, 1998, "Quantitative, competitive PCR assay for HIV-1 using a 30 microplate-based detection system", Biotechniques 24:810-6); 11) colorimetric procedures; and 12) biological assays (e.g., for virus titers).

In a particular embodiment, protein-probe interactions are detected by direct mass spectrometry. In a further embodiment, the identity of the protein and/or probe is determined using mass spectrometry. For example, one of more probes that have bound to a protein on the proteome chip can be dissociated from the array, and identified by mass

spectrometry (see, e.g., WO 98/59361). In another example, enzymatic cleavage of a protein on the proteome chip can be detected, and the cleaved protein fragments or other released compounds can be identified by mass spectrometry.

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In one embodiment, each protein on the proteome chip is contacted with a probe, and the protein-probe interactions are detected and quantified. In another embodiment, each protein on the proteome chip is contacted with multiple probes, and the protein-probe interaction is detected and quantified. For example, the proteome chip can be simultaneously screened with multiple probes including, but not limited to, complex mixtures (e.g., cell extracts), intact cellular components (e.g., organelles), whole cells, and 10 probes pooled from several sources. The protein-probe interactions are then detected and quantified. Useful information can be obtained from assays using mixtures of probes due, in part, to the positionally addressable nature of the arrays of the present invention, i.e., via the placement of proteins at known positions on the protein chip, the protein to which the probe binds ("interactor") can be characterized.

15 One of ordinary skill in the art can appreciate many different embodiments for assaying various cellular interactions by using probes to screen the proteome chips of the invention. For example, multiple sequential screens of a proteome chip with various probes can define all proteins involved in a particular signal transduction pathway or in a specific metabolic pathway. Moreover, these assays can be useful for diagnostic, prognostic and/or 20 therapeutic purposes.

In accordance with the methods of the invention, a probe can be a cell, cell membrane, subcellular organelles, protein-containing cellular material, protein, oligonucleotide, polynucleotide, DNA, RNA, small molecule (i.e., a compound with a molecular weight of less than 500), substrate, drug or drug candidate, receptor, antigen, 25 steroid, phospholipid, antibody, immunoglobulin domain, glutathione, maltose, nickel, dihydrotrypsin, lectin, or biotin.

Probes can be biotinylated for use in contacting a protein array so as to detect protein-probe interactions. Weakly biotinylated proteins are more likely to maintain the biological activity of interest. Thus, a gentler biotinylation procedure is preferred so as to 30 preserve the protein's binding activity or other biological activity of interest. Accordingly, in a particular embodiment, probe proteins are biotinylated to differing degrees using a biotin-transferring compound (e.g., Sulfo-NHS-LC-LC-Biotin; PIERCETM Cat. No. 21338, USA).

In addition, the probe can be an enzyme substrate or inhibitor. For example, the 35 probe can be a substrate or inhibitor of an enzyme such as, but not limited to, kinases,

phosphatases, proteases, glycosidases, acetylases, and other group transferring enzymes. After incubation of proteins on a chip with combinations of nucleic acid or protein probes, the bound nucleic acid or protein probes can be identified, for example, by mass spectrometry (Lakey et al., 1998, "Measuring protein-protein interactions", Curr Opin Struct Biol. 8:119-23).

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Accordingly, various cellular responses to interaction with the proteins on a proteome chip can be assayed by probing with whole cells. For example, a proteome chip can be contacted with lymphocytes and assayed for lymphocyte activation by various means including, but not limited to, detecting antibody synthesis, detecting or measuring incorporation of ³H-thymidine, labeling cell surface molecules with antibodies to identify molecules induced or suppressed by antigen recognition and activation (*e.g.*, CD23, CD38, IgD, C3b receptor, IL-2 receptor, transferrin receptor, membrane class II MHC molecules, PCA-1 molecules, HLA-DR), and identifying expressed and/or secreted cytokines.

In another example, mitogens for a specific cell-type can be determined by

incubating a cell with a proteome chip. Mitotic activity can be determined, for example, by
detecting or measuring incorporation of ³H-thymidine by a cell. Cells can be of the same
cell type (*i.e.*, a homogeneous population) or can be of different cell types.

In another example, differentiation factors for a specific cell-type can be determined by incubating a cell with a proteome chip. Differentiation of a cell can be determined, for example, by visual inspection, detection of cell-surface differentiation markers using marker-specific antibodies, or identification of secreted differentiation markers.

In another example, apoptotic factors for a specific cell-type can be determined by incubating a cell with a proteome chip. Apoptosis can be assayed, for example, by visual inspection, detection of cell-surface apoptotic markers using marker-specific antibodies, or identification of secreted markers or other cellular components released into the media.

In another example, the secretory response of a cell to a protein on a proteome chip can be assayed by incubating a cell with a proteome chip of the invention. Secreted proteins and other cellular compounds can be assayed, for example, by detecting the released compounds in the media.

In another example, the ability of a protein on a proteome chip to mediate cell aggregation can be assayed, for example, by incubating one or more cells with a proteome chip of the invention, and assaying for aggregation. Also, a protein's ability to mediate an affinity to extracellular matrix can be assayed by, for example, incubating a cell and extracellular matrix components with a proteome chip, and assaying for enhanced affinity of the cell or the extracellular matrix component with a protein on the chip. Interactors

identified in such assays can have a role in, for example, cancer, cell migration, synaptogenesis, dendritic growth, process extension, or axonal elongation.

In yet another example, the effect of proteins of a proteome chip of the invention on ion transport, or other small molecule transport (e.g., ATP), can be determined. For example, the probe cells can be pre-loaded with a radioactively labeled ion or other small molecule, and incubated with a proteome chip of the invention. Retention or release of the radioactive label can be measured at different time points after contacting the cells with the proteins of the proteome array. Alternatively, ion transport can be detected and characterized using electrophysiological techniques known in the art.

In yet another example, cellular uptake and/or processing of proteins on the proteome chips can be assayed by, for example, incubating a cell with a proteome chip having radioactively or fluorescently labeled proteins on the chip, and measuring the increase or decrease in signal on the proteome chip, or measuring uptake of labeled protein by the cell.

Alternatively, a proteome chip of the invention can be incubated with a cell and a labeled compound of interest, such that cellular uptake and/or processing of the compound by the cell is detected and/or measured.

Interactions of small molecules (*i.e.*, compounds smaller than MW=500) with the proteins on a proteome chip also can be assayed in a cell-free system by probing with small molecules such as, but not limited to, ATP, GTP, cAMP, phosphotyrosine, phosphoserine, and phosphothreonine. Such assays can identify all proteins in a species that interact with a small molecule of interest. Small molecules of interest can include, but are not limited to, pharmaceuticals, drug candidates, fungicides, herbicides, pesticides, carcinogens, and pollutants. Small molecules used as probes in accordance with the methods of the invention preferably are non-protein, organic compounds.

In another embodiment, essentially all receptors for a particular ligand, or class of ligands, in a species can be identified by contacting a receptor of interest with a proteome chip of the invention. Alternatively, essentially all ligands in a species that are identified by a particular receptor or receptor family of interest can be identified by contacting a receptor of interest with a proteome chip of the invention. In another embodiment, essentially all proteins in a species, capable of inhibiting or blocking formation of a particular receptor-ligand complex, can be identified by contacting a receptor and its ligand with a proteome chip of the invention, and determining whether receptor-ligand interaction is inhibited as compared with the degree of receptor-ligand interaction in the absence of the

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protein on the chip. Detection of receptor-ligand interaction and identification of the ligand interactors can be accomplished using methods known in the art.

In another embodiment, essentially all kinase targets in a species can be identified by, for example, contacting a kinase with a proteome chip of the invention, and in the presence of labeled phosphate, detecting phosphorylated interactors using methods known in the art. Alternatively, essentially all kinases in a species can be identified by contacting a substrate that can be phosphorylated with a proteome chip of the invention, and assaying the presence and/or level of phosphorylated substrate by, for example, using an antibody specific to a phosphorylated amino acid. In another embodiment, essentially all kinase inhibitors in a species can be identified by contacting a kinase and its substrate with a proteome chip of the invention, and determining whether phosphorylation of the substrate is reduced as compared with the level of phosphorylation in the absence of the protein on the chip.

Detection methods for kinase activity are known in the art, and include, but are not limited to, the use of radioactive labels (e.g., ³³P-ATP and ³⁵S-γ-ATP) or fluorescent antibody probes that bind to phosphoamino acids.

Similarly, assays can be conducted to identify all phosphatases, and inhibitors of a phosphatase, in a species. For example, whereas incorporation into a protein of radioactively labeled phosphorus indicates kinase activity in one assay, another assay can be used to measure the release of radioactively labeled phosphorus into the media, indicating phosphatase activity.

The proteome chips of the invention can also be used to distinguish different cell types (either morphological or functional) by, for example, contacting a proteome chip with cells or cell extracts representing different populations of cells, and comparing the patterns of protein-probe interactions on the proteome chip. This approach also can be used to characterize, for example, different stages of the cell cycle, disease states, altered physiologic states (e.g., hypoxia), physiological state before or after treatment (e.g., drug treatment), metabolic state, stage of differentiation, developmental stage, response to environmental stimuli (e.g., light, heat), response to environmental toxins (e.g., pesticides, herbicides, pollution), cell-cell interactions, cell-specific protein expression, and disease-specific protein expression.

Developmental profiles of protein-protein interactions can be used to characterize signal transduction pathways, metabolic pathways, etc. involved at every development stage and elucidate transitions between developmental stages. The wealth of information

provided by such studies can be used to identify drug targets for each stage, and/or tailor treatment regimens during the course of a disease.

The proteome chips of the invention can be incubated with cell extracts to characterize a particular cell type, response to a stimulus, or physiological state.

Accordingly, in exemplary embodiments, a proteome chip of the invention can be contacted with a cell extract from cells treated with a compound (e.g., a drug), or from cells at a particular stage of cell differentiation (e.g., pluripotent), or from cells in a particular metabolic state (e.g., mitotic), and assayed for kinase, protease, glycosidase, actetylase, phosphatase, and/or other transferase activity, for example.

The pattern of protein-probe interactions on the proteome chip can thereby provide a "signature" or "fingerprint" characteristic of the biological state. For example, the results obtained from such assays, comparing for example, cells in the presence or absence of a drug, or cells at several differentiation stages, or cells in different metabolic states, can provide a signature of each condition, and can provide information regarding the physiologic changes in the cells under the different conditions.

Clearly, by screening a species's proteome using a plurality of probes (*e.g.*, known mixtures of probes, cellular extracts, subcellular organelles, cell membrane preparations, whole cells, etc.), the resulting analysis of protein-probe interactions can form the basis of identifying a "fingerprint" or "signature" of the a cell-type or physiological state of a cell, tissue, organ or system. Such information can be useful for diagnosis, prognosis, drug testing, and drug discovery, for example.

Accordingly, the proteome chips of the invention can be used to determine a drug's interactions with proteins on the chip. Alternatively, the proteome chips of the invention can be used to characterize a drug's effects on complex protein mixtures such as, for example, whole cells, cell extracts, or tissue homogenates. For example, a proteome chip can be contacted with a complex protein mixture and assayed for altered interactions of the protein mixture with the proteins on the chip when compared in presence or absence of drug.

The net effect of a drug can thereby be analyzed by screening one or more proteome 30 chips with drug-treated cells, tissues, or extracts, which then can provide a "signature" for the drug-treated state, and when compared with the "signature" of the untreated state, can be of predictive value with respect to, for example, potency, toxicity, and side effects. Furthermore, time-dependent effects of a drug can be assayed by, for example, adding the drug to the cell, cell extract, tissue homogenate, or whole organism, and applying the

drug-treated cells or extracts, prepared at various timepoints of the treatment, to a proteome chip. Such assays can be useful for diagnosis or prognosis of a disease.

In particular, the proteome chips of the invention can be useful for characterizing a mode of action of a drug, determining drug specificity, predicting drug toxicity, and for drug discovery. For example, the identity of proteins that bind to a drug, and their relative affinities, can be assayed by incubating a proteome chip with a drug or drug candidate under different assay conditions, determining drug specificity by determining where on the array the drug bound, and measuring the amount of drug bound by each different protein.

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The proteome chips of the invention can be used to determine a disease state by, for example, contacting a proteome chip with diseased cells, cell extracts or tissue homogenates from diseased tissue, or body fluids from a patient suffering from a disease, and comparing the pattern of protein-probe interactions on the proteome chip with that of a healthy counterpart. Such assays can provide a "signature" for the disease state, and when compared with the "signature" of the healthy state, can be of predictive value with respect to, diagnosis or prognosis of the disease. Furthermore, stages of a disease can be characterized by, for example, assaying biological preparations on the proteome chip at various stages of the disease.

Bioassays in which a biological activity is assayed, rather than binding assays, can also be conducted out on the same proteome chip, or on an identical second chip. Thus, these types of assays using the protein chips of the invention are useful for studying drug specificity, predicting potential side effects of drugs, and classifying drugs.

Further, proteome chips of the invention are suitable for screening complex libraries of drug candidates. Specifically, the proteins on the chip can be incubated with the library of drug candidates, and then the bound components can be identified, *e.g.*, by mass spectrometry, which allows for the simultaneous identification of all library components that bind preferentially to specific subsets of proteins, or bind to several of the proteins on the chip. Additionally, the relative affinity of the drug candidates for the different proteins in the array can be determined.

Moreover, the protein chips of the present invention can be probed in the presence of potential inhibitors, catalysts, modulators, or enhancers of an observed interaction, enzymatic activity, or biological response. Using a proteome chip of the present invention, such strategic screens can identify proteins expressed in a species that, for example, block the binding of a drug, inhibit of viral infection, exhibit bacteriostatic activity, exhibit anti-fungal activity, ameliorate parasitic infection, or physiological effectors to specific categories of proteins.

Enzymatic reactions can be performed and enzymatic activity measured using the proteome chips of the present invention. In a specific embodiment, compounds that modulate the enzymatic activity of a protein or proteins on a chip can be identified. For example, changes in the level of enzymatic activity can be detected and quantified by incubating a compound or mixture of compounds with an enzymatic reaction mixture, thereby producing a signal (e.g., from substrate that becomes fluorescent upon enzymatic activity). Differences between the presence and absence of a test compound can be characterized. Furthermore, the differences in a compound's effect on enzymatic activities can be detected by comparing their relative effect on samples within the proteome chip and between chips.

A variety of strategies of using the proteome chips of the present invention can be employed to determine various physical and functional characteristics of proteins. For example, the protein chips can be used to assess the presence and amount of protein present by probing with an antibody. The protein can be detected using standard detection assays such as luminescence, chemiluminescence, fluorescence, chemifluorescence, or mass spectrometry. For example, a primary antibody to the protein of interest is recognized by a fluorescently labeled secondary antibody, which is then measured with an instrument (e.g., a Molecular Dynamics scanner) that excites the fluorescent product with a light source and detects the subsequent fluorescence. For greater sensitivity, a primary antibody to the protein of interest is recognized by a secondary antibody that is conjugated to an enzyme such as alkaline phosphatase or horseradish peroxidase. In the presence of a luminescent substrate (for chemiluminescence) or a fluorogenic substrate (for chemifluorescence), enzymatic cleavage yields a highly luminescent or fluorescent product which can be detected and quantified by using, for example, a Molecular Dynamics scanner.

Alternatively, the signal of a fluorescently labeled secondary antibody can be amplified using an alkaline phosphatase-conjugated or horseradish peroxidase-conjugated tertiary antibody.

In one embodiment, a proteome chip of the invention can probed with antibodies directed against known proteins in one species, such that homologous proteins having recognized epitopes can be identified in the proteome of another species. Same species homologs of the interactors can be obtained by, for example, using the DNA sequence information of the homologous protein to identify the homolog in the other species. In specific embodiments, the antibody is directed against cyclin, kinase, GST, Clb5, Cla4, Ste20, Cdc42, PI(3,4)P2, PI(4)P, SPA2, CLB1, CLB2, or Cdc11.

In another embodiment, a proteome chip of the invention containing proteins of a first species can probed with a protein from a second species to identify interactors. Homologs in the second species of the interactors from the first species can then be identified and characterized by, for example, nucleotide sequence homology. Thus, where the proteome is not available for a particular species, this strategy can be used to find same species proteins that interact with a protein of interest.

The proteome chips of the invention can be used to identify essentially all substrates in a species for each enzyme found in a species. Accordingly, identifying substrates of protein kinases, phosphatases, proteases, glycosidases, acetylases, or other group transferring enzymes can be conducted on the protein chips of the present invention.

In one embodiment, protease activity can be detected by identifying, using standard assays (*e.g.*, mass spectrometry, fluorescently labeled antibodies to peptide fragments, or loss of fluorescence signal from a fluorescently tagged substrate), peptide fragments that are produced by protease activity and released into the media. Thus, activity of group-transferring enzymes can be assayed readily by several approaches using any means of detection, which would be appreciated by one of ordinary skill in the art.

The proteome chips of the invention can be used to identify essentially all binding proteins in a species for any compound. Accordingly, proteome chips can be used to identify and characterize essentially all proteins that bind, for example, kinases, proteases, hormones, DNA, RNA, phosphatases, proteases, glycosidases, acetylases, or other group transferring enzymes. Thus, the chip can be probed with a probe, and assayed for protein-probe interaction and/or assayed for the desired activity. For example, if RNA binding is the activity of interest, the proteome chip is probed with RNA, and protein-RNA complexes are identified.

25 For example, the proteome chips of the invention can be used to identify essentially all proteins in a species that bind to membrane-associated proteins or other membrane-associated compounds by contacting the chip with probes such as, for example, whole cells, preparations of plasma membranes, membrane vesicles, or liposome comprising membrane components of interest, and detecting protein-probe interaction. In a particular embodiment, the probe is in the form of a liposome comprising one or more phospholipids of interest. The protein-probe interaction can be detected using techniques known in the art. The identity of the interactor and/or probe can be determined using techniques known in the art. Moreover, biological activities (e.g., enzyme activity, cell activation) can also be detected using techniques known in the art.

The identity of target proteins from pathogens (e.g., an infectious disease agent such as a virus, bacterium, fungus, or parasite) or target proteins from abnormal cells (e.g., neoplastic cells, diseased cells, or damaged cells) that serve as antigens in the immune response of recovering or non-recovering patients can be determined by using a proteome chip of the invention. For example, lymphocytes isolated from a patient can be used to screen chips comprising all of a pathogen's proteins. In general, these screens comprise contacting a proteome chip with a plurality of lymphocytes, wherein the proteins on the proteome chip comprise a plurality of potential antigens, and detecting positions on the chip where lymphocyte activation occurs. In a specific embodiment, lymphocytes are contacted 10 with a pathogen's proteins on the proteome chip, after which activation of B-cells or T-cells by an antigen or a mixture of antigens is assayed, thereby identifying target antigens derived from a pathogen.

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Alternatively, the proteome chips of the invention can be used to characterize an immune response by, for example, screening a proteome of an infectious organism with a 15 patient's lymphocytes to identify the targets of a patient's B-cells and/or T-cells. For example, B-cells can be incubated with a proteome chip of the invention to identify antigenic targets for humoral-based immunity.

In another embodiment, the proteome chips of the invention can be used to detect and characterize substrates of autoimmunity or allergy-causing proteins. For example, a 20 proteome of human proteins can be screened, with a patient's lymphocytes or with a patient's circulating antibodies, to identify the targets of a patient's B-cells and/or T-cells. Such screens can characterize autoimmunity or allergic reactions, and identify potential target drug candidates.

In one embodiment, the proteome chips of the invention are used to identify 25 substances that are able to activate B-cells or T-cells. For example, lymphocytes are contacted with the proteome chip, and lymphocyte activation is assayed, thereby identifying substances that have a general ability to activate B-cells or T-cells or subpopulations of lymphocytes (e.g., cytotoxic T-cells).

Induction of B-cell activation by antigen recognition can be assayed by various 30 means including, but not limited to, detecting or measuring antibody synthesis, incorporation of ³H-thymidine, binding of labeled antibodies to newly expressed or suppressed cell surface molecules, and secretion of factors indicative of B-cell activation (e.g., cytokines). Similarly, T-cell activation in a screen using a protein chip of the invention can be determined by various assays. For example, a chromium (51Cr) release 35 assay can detect recognition of antigen and subsequent activation of cytotoxic T-cells (see,

e.g., Palladino et al., 1987, Cancer Res. 47:5074-9; Blachere et al., 1993, J. Immunotherapy 14:352-6).

The specificity of an antibody preparation can be determined through the use of a proteome chip of the invention, comprising contacting the chip with an antibody

5 preparation, and detecting positions on the solid support where binding by an antibody in the antibody preparation occurs. The antibody preparation can be, but is not limited to, Fab fragments, antiserum, and polyclonal, monoclonal, chimeric, single chain, humanized, or synthetic antibodies. In one example, a proteome chip is probed with a monoclonal antibody to characterize its binding strength and/or its specificity. In specific embodiments, the antibody is directed against cyclin, kinase, GST, Clb5, Cla4, Ste20, Cdc42, PI(3,4)P2, PI(4)P, SPA2, CLB1, CLB2, or Cdc11.

The proteome chips of the invention are useful for identifying proteins that bind to specific molecules of biologic interest including, but not limited to, receptors for potential ligand molecules, virus receptors, and ligands for orphan receptors.

The proteome chips are also useful for detecting DNA binding or RNA binding to proteins on the chips, and for evaluating the binding specificity and strength. The DNA can be single-stranded or double-stranded. The RNA can be mRNA, hnRNA, polyA⁺ RNA, or total RNA.

The proteome chips of the invention are useful for identifying proteins that are modified posttranslationally. Posttranslational modifications that can be detected using the methods of the invention include, but are not limited to, methylation, acetylation, farnesylation, biotinylation, stearoylation, formylation, lipoic acid, myristoylation, palmitoylation, geranylgeranylation, pegylation, phosphorylation, sulphation, glycosylation, sugar modification, lipidation, lipid modification, ubiquitination, sumolation, disulphide bonding, cysteinylation, oxidation, glutathionylation, pyroglutamic acid, carboxylation, and

deamidation. In addition, a protein can be modified posttranslationally with pentoses, hexosamines, N-acetylhexosamines, deoxyhexoses, hexoses, and sialic acid.

In one embodiment, a proteome chip of the invention can be probed with phosphotyrosine, phosphoserine, or phosphothreonine to identify phosphorylated interactors. In another example, a proteome chip of the invention can be probed with a lectin (e.g., wheat germ agglutinin) to identify glycosylated interactors (e.g., N-acetylglucosamine). The phosphorylated or glycosylated interactors can be detected using methods known in the art.

The proteome chips of the invention are also useful for identifying and characterizing protein isoforms that exhibit differences in function, ligand binding, or

enzymatic activity. In a particular embodiment, a proteome chip of the invention is used to characterize different binding affinities by protein isoforms derived from different alleles by assaying their activities relative to one another.

In one embodiment, consensus sequences can be determined using the proteome chips of the invention. For example, upon screening a proteome chip with a binder (which can be from a species different from that of the proteins on the chip), the amino acid sequences of the interactors can be aligned to construct a consensus sequence for the interacting domain(s). Such information can be useful, for example, for designing inhibitors of the binder-interactor interaction, or for designing novel and/or improved 10 binders for a particular interactor or class of interactors.

In a specific embodiment, a consensus sequence for calmodulin-binding proteins is determined by screening a proteome chip with calmodulin. In a further embodiment, the consensus sequence for a calmodulin-binding protein consists of the amino acid sequence I/L-Q-X-K-K/X-G-B (SEQ ID NO: 1). In another embodiment, the consensus sequence for 15 a calmodulin-binding protein comprises the amino acid sequence I/L-Q-X-K-K/X-G-B (SEQ ID NO: 1). In a further embodiment, the consensus sequence for a calmodulin-binding protein comprises the amino acid sequence I/L-Q-X-K-K/X-G-B (SEQ ID NO: 1), and is less than 10, 15, 30, 50, or 100 amino acid residues in length.

The proteome chips of the invention are also useful for identifying a set of potential 20 antibacterial, antifungal, antiparasitic or antiviral compounds. For example, cell lysates or other preparations of bacteria, fungi, parasites or viruses can be contacted with a proteome array, and protein-probe interactions detected and identified. Additionally, comparing interaction patterns obtained from infectious organisms at infectious stages and non-infectious stages can identify interactions involved in infection of the host.

25 Screening of phage display libraries can be performed by incubating a library with the proteome chips of the present invention. The detection of clones binding to a protein on the chip can be conducted by various methods known in the art (e.g., mass spectrometry), thereby identifying clones of interest, after which the DNA encoding the clones of interest can be identified by standard methods (see, e.g., Ames et al., 1995, J. Immunol. Methods 30 184:177-86; Kettleborough et al., 1994, Eur. J. Immunol. 24:952-8; Persic et al., 1997, Gene 187:9-18). In this manner, the chips are useful to select for cells having surface components that bind to specific proteins on the chip.

5.5 Kits.

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The invention also provides kits for carrying out the assay regimens of the invention. In one embodiment, kits of the invention comprise one or more proteome chips of the invention. Such kits may further comprise, in one or more containers, reagents useful for assaying biological activity of a protein or molecule, reagents useful for assaying protein-probe interaction, and/or one or more probes, proteins or other molecules. The reagents useful for assaying biological activity of a protein or other molecule, or assaying interactions between a probe and a protein or other molecule, can be applied with the probe, attached to a proteome chip, or contained in one or more wells on a proteome chip. Such 10 reagents can be in solution or in solid form. The reagents may include either or both the proteins or other molecules and the probes required to perform the assay of interest.

The proteins of the proteome chip can be attached to the surface of a flat solid support, contained in wells on a solid support, or attached to the surface of wells on the solid support. In one embodiment, the proteome chip in the kit has the proteins and/or probes already attached to the solid support. In another embodiment, the proteome chip in the kit can have the reagent(s) or reaction mixture useful for assaying biological activity of a protein or other molecule, or useful for assaying the interaction of a probe and a protein or other molecule, already attached to wells on the solid support. In yet another embodiment, the reagent(s) is not attached to the wells of the solid support, but is contained in the wells. 20 In yet another embodiment, the reagent(s) is not attached to the wells of the solid support, but is contained in one or more containers, and can be added to the wells of the solid support. In yet another embodiment, the kit further comprises one or more containers holding a solution reaction mixture for assaying biological activity of a protein or molecule. In yet another embodiment, the kit provides a substrate (e.g., beads) to which probes, 25 proteins or molecules of interest, and/or other reagents useful for carrying out one or more assays, can be attached, after which the substrate with attached probes, proteins, or other reagents can be placed into the wells of the chip.

5.6 Design of a Positive Identification Algorithm.

30 The present invention also provides a method for identifying whether a signal is positive. Signals can be in any measurable form including, but not limited to, visible light, ultraviolet radiation, infrared radiation, X-rays, fluorescence, and colorimetric visualization.

In a preferred embodiment, signals are detected by mass spectrometry.

In addition, signals can be in any arrangement, and in any physical format such as, but not limited to, arrays, blots, gels, and screens. Preferably, signals are spots in a static arrangement.

In another preferred embodiment, signals are produced by fluorescence and are arranged in a grid pattern on an array. As such, a signal can be assigned a positional coordinate with respect to row and column. The rows and columns can be of any width.

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The first step in filtering signals is to calculate the local foreground and background signals for each spot. The local foreground signal is emitted from the actual spot, whereas the local background signal is emitted from the area immediately bordering to the spot. The net signal, which is the local foreground signal minus the local background signal, is used in all further calculations. The local foreground and background signals can be identified by software such as GENEPIXTM.

However, variations between chips, which can represent, for example, different lipid-binding experiments, and local variations on the chip (due to unequal diffusion of substrates, for example) can result in further fluctuation of the net signal intensity, resulting in different net signal distributions for different chips. To correct the variation between chips, the net signals from different chips need to be scaled into a common range. One of the several chips is chosen as a reference and the goal is to scale the net signal distributions of each chip to the range and shape of the net signal distribution of the reference chip.

For example, the lower quartile, median, and upper quartile values of the net signal distribution of each chip can be computed. Then, for each chip, the median net signal is subtracted from the net signal of each spot. Furthermore a scaling factor is computed for each chip, which is equal to the ratio of the difference between the upper and lower quartile of the specific chip and the difference between the upper and lower quartile of the reference chip. This implies that the scaling factor for the reference chip is equal to one. Then the net signals on each chip are multiplied by the chip-specific scaling factor to calculate the scaled net signals.

To correct for local variations on an array, a "neighborhood subtraction" for each spot can be performed. For example, the neighborhood region can be defined as a region of two rows above and below, as well as two columns to the left and right of a signal spot. The median signal of this region is then subtracted from the spot signal to calculate an excess signal relative to the neighborhood of the spot. Preferably, the number of spots of high signal strength in any neighborhood region is sufficient low, such that the median value is not significantly affected and is a good representation of the background signal in the neighborhood region.

Applying the neighborhood subtraction to the scaled net signals yields the scaled excess signals. In the next step, parallel samples are compared with respect to their scaled excess signals. If the difference between the average of the scaled excess signal of the two parallel samples and the scaled excess signal of one of the parallel samples is greater than three times the standard deviation of the error of the scaled excess signal, the spots belonging to the two parallel samples are excluded from further analysis. The remaining spots and their scaled excess signal then represent the set of filtered signals.

The distribution of the error of the scaled excess signals and its standard deviation can be computed as follows. A linear regression is performed on the complete set of parallel samples to determine the general linear relationship between parallel samples. Then an error value can be calculated for each parallel sample:

$$\varepsilon_G = |G2 - Gm|,$$

where

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G1 and G2 represent the two scaled excess signals of the two parallel samples,

15 Gm = (G1 + G2)/2,

and function f(G1) = a*G2 + b is the general linear relationship between the two parallel samples G1 and G2, with the parameters a and b determined by linear regression.

The complete set of error values is the set of error values for all parallel samples and represents the distribution of error values for the scaled excess signal. Then the standard deviation of this distribution can be calculated.

Finally, a pair of parallel samples is called positive if the average of their filtered signals is three standard deviations greater than the error of the scaled excess values. (Note that this threshold is independent of the threshold to determine whether parallel samples should be excluded from the set of filtered signals.)

After this filtering procedure, the filtered signal (G) is normalized with the GST signal (R), yielding the ratio r = G/R which can be a measure of the binding per amount of protein and can allow for comparison of binding signals between different proteins. The specific binding ratio r is sensitive to errors ε_G and ε_R in both the G and R signals. Using a Monte-Carlo procedure, 90% and 95% confidence intervals for this ratio can be calculated.

The error e_G of the r value is related to the errors e_G and e_R :

$$r + \varepsilon_R = (G + \varepsilon_G)/(R + \varepsilon_R)$$

where ε_R represents the error of the ratio r.

For the Monte Carlo procedure, both the distributions of ε_G and ε_R must be known. The distribution of e_G can be computed as explained above. The distribution of e_R can be

computed in the same procedure as for e_G by using the GST signal pairs of parallel samples as input.

The Monte Carlo procedure is as follows:

In order to determine confidence intervals for $r + \varepsilon_R$, a population of random samples of $r + \varepsilon_R$ needs to be computed. These can be derived from random samples of ε_G and ε_R . Random samples of ε_G can be computed as follows. Samples of uniformly distributed random numbers between 0 and 1 are calculated with a standard random number generator. From the distributions of ε_G the inverse cumulative distribution function of ε_G is determined by standard procedures. Using the samples of random numbers as arguments for this function produces a set of random samples for ε_G . Likewise a set of random samples for ε_G can be calculated. These random samples of ε_G and ε_R can be combined in the formula $r + \varepsilon_R = (G + \varepsilon_G)/(R + \varepsilon_R)$ to produce a population of random samples for $r + \varepsilon_R$.

In one embodiment, the invention provides a method for identifying whether a signal is positive, comprising the steps of determining foreground and background signals for each spot locally and determining net signals from their difference, determining the lower quartile, median, and upper quartile values of a first and second net signal distribution, subtracting the first median value from the first net signal distribution, and subtracting the second median value from the second net signal distribution to obtain a first and second subtracted value, respectively, dividing said first subtracted value by the difference between said upper and lower quartile values of said first signal distribution, and dividing said second scaled value by the difference between said upper and lower quartile values to obtain a first and second scaled value, respectively, computing a local median value of a scaled signal distribution of a neighborhood region, wherein said neighborhood region comprises a plurality of sites in the area; subtracting the local median value from the scaled signal to obtain an scaled excess value; and parallel samples of scaled excess values are excluded if the difference between one of the sample values and their average is greater than three standard deviations of the error of the scaled excess value.

The filtered values can be used to identify whether a signal is positive. Parallel samples are called "positive" if the average of the filtered values of the parallel sample is three times greater than the standard deviation of the error of scaled excess values. Filtered positive signals can then be normalized using the formula:

r = G/R

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wherein G is the filtered value, R is a GST signal, and r represents a signal per amount of protein that can be compared among different spots. The ratio r is sensitive to the errors in G and R. This sensitivity can be assessed by calculating confidence intervals of

$$r + \varepsilon_r = G + \varepsilon_G / R + \varepsilon_R$$

 ε_G is the error of G, ε_R is the error of R, and ε_r is the error of r.

In a specific embodiment, a positive signal indicates protein-probe interaction.

In another specific embodiment, the neighborhood region is two rows above, two rows below, two columns to the left, and two columns to the right of the signal.

In another specific embodiment, data points from two parallel samples are excluded from further analysis, wherein the difference between the scaled excess signals of said samples and their average is greater than three standard deviations of the error of the scaled excess signal. Data points are also excluded if they are obviously artifactual.

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6. EXAMPLES

A defined collection of over 5800 proteins from the budding yeast was prepared using high-throughput techniques and screened for many activities including protein-protein, protein-DNA, protein-RNA, and protein-liposome interactions. A large number of novel activities were identified, providing new information concerning known and previously uncharacterized genes.

To facilitate studies of the yeast proteome, 5800 open reading frames were cloned and overexpressed, and their corresponding proteins purified. The proteins were printed onto slides at high spatial density to form a yeast proteome microarray and screened for their ability to interact with proteins and phospholipids. Many novel calmodulin and phospholipid-interacting proteins were identified; a common potential binding motif was identified for many of the calmodulin-binding proteins. These studies demonstrate that microarrays of an entire eukaryotic proteome can be prepared and screened for large numbers of biochemical activities resulting in the identification of many novel protein functions/interactions. These microarrays can also be screened to detect protein posttranslational modifications.

6.1 Yeast Culture Preparation

- Yeast glycerol stocks stored in 96-well plates at -80°C were inoculated onto a URA-agar plate (Omni, USA) using a 96-pronger.
 - 2. The culture was allowed to grow on agar at 30°C for 48 hours, and visible colonies (2 mm diameter) can be observed.

3. A 96-pronger was used to inoculate yeast cells from agar plates to a 96-well 2 ml box in which every well contained 300 µl URA-/raffinose liquid media and a 2 mm diameter glass ball, which facilitates the uniform growth.

- After the culture reached O.D.₆₀₀ 4.0 in about 16 hours at 30°C with vigorous
 shaking (300 rpm), 15 μl of the same strain was inoculated into 750 μl of
 URA-/raffinose liquid media in four different boxes to obtain 3 ml of culture.
 Again, each well contained the same glass ball to achieve aeration and even growth.
 The cells were grown at 30°C with vigorous shaking.
- 5. After 12-16 hours of growth, the culture should reach O.D.₆₀₀ 0.6 to 0.8. Cultures were discarded if the OD is over 1.0. Using an automated liquid-handling device (Q-Fill, Genetix, UK), 40% galactose stock was added to each well to a final concentration of 2% to induce the cells. The cultures were induced at 30°C for 4 hours with shaking.
- The cells were harvested by spinning at 3000 rpm for 2-10 min, and the cell pellets were resuspended in 100-1000 μl of cold water by vortexing. Cells of the same strain were then merged from 4 wells into one. Cells were collected by spinning and resuspended in 100-1500 μl of cold lysis buffer without the protease inhibitors on ice. The washed cells were collected by a brief centrifugation, and the lysis buffer was discarded. The washed semi-dry culture was immediately stored in -80°C
 freezer. The culture can be kept for weeks.

6.2 Protein Purification in a 96-well Fashion

- The frozen culture in a 96-well box was transferred from -80°C to ice and 100-300 μl of zirconia beads (0.5 mm diameter from BSP, Germany) was added to each well. While the culture was still frozen, 100-500 μl of lysis buffer containing fresh protease inhibitors was added. A cap mat was used to seal each well. After thawing the culture for 5-25 min on ice, the cells in the 96-well box were vortexed 20-60 seconds for 3-6 times with 1-5 min intervals on ice. To efficiently disrupt the yeast cell wall, and to process many plates at once, a paint shaker (HARBILTM 5G HD, 36 kg capacity, adjustable pressure and shaking time, fixed speed at 200 times per minute) was used to violently agitate the samples.
- After spinning at 3000 rpm for 2-10 min, the supernatant was collected using wide-open tips (Fisher, USA) and transferred into a 96-well filter plate (Whatman, USA; Whatman UNIFILTERTM, Cat. No. 7700-1801 having a hydrophilic PVDF filter with an 800 μl/well capacity), which was placed on top of a 96-well box.

3. To obtain more proteins, 100-500 µl of lysis buffer containing fresh protease inhibitors was added to the cell debris, and Steps 1 and 2 were repeated.

- 4. The combined cell lysate was spun through the filter plate into a cold and clean 96-well box for 10-30 min at 3000 rpm. The volume of filtered lysate in each well was roughly 200-1000 μl.
- 5. Meanwhile, the required amount of glutathione beads (roughly 10-50 μl of beads per sample) (Amersham, USA) was washed four times with cold lysis buffer without the protease inhibitors, and finally resuspended in 5X of its original volume with lysis buffer containing fresh protease inhibitors.
- 100 μl of washed glutathione beads was added to each well and sealed tightly with a cap mat. The beads were incubated with the lysate on a roller drum at 4°C for one hour. To obtain the best mixing, the boxes were rotated 360 degrees on the roller drum.
- 7. The beads were collected by spinning at 3000 rpm for 10-60 seconds, and the supernatant was discarded. Beads were washed once with 200-800 μl of Wash Buffer I containing protease inhibitors, and twice without the inhibitors.
- 8. The beads were then washed three times with 200-800 μl of Wash Buffer II. After complete removal of the buffer, 20-50 μl of Elution Buffer was added to each well. Filter plates used for the elution step comprised materials having low affinity for proteins (MILLIPORE MULTISCREENTM, Cat. No. MADVN6550 having a hydrophilic PVDF filter with a 200 μl/well capacity). The box was vortexed briefly to resuspend the beads and incubated on a roll drum for one hour at 4°C.
- 9. The eluate/beads slurry was transferred to a cold filter plate (Millipore, USA), and the eluate was collected to a 96-well PCR plate by spinning through the filter plate for 0.5-2 min at 3000 rpm at 4°C.
 - 10. Each purified protein was aliquoted into three 96-well PCR plates and immediately stored in a -80°C freezer.

30 Lysis Buffer

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30-300 mM Tris pH 7.5
50-300 mM NaCl
0.1-10mM EGTA
0.01-1.0% TritonX-100

0.01-1% beta-mercaptoethanol ("BME")
0.1-3 mM phenylmethylsulfonyl fluoride ("PMSF")

Roche Protease inhibitor tablets (containing EDTA) BME, PMSF, and inhibitor tablets were added freshly.

Wash Buffer I:

5 30-300 mM Tris pH 7.5 300-600 mM NaCl 0.1-10mM EGTA 0.01-1.0% TritonX-100

0.01-1% beta-mercaptoethanol ("BME")

10 0.1-3 mM PMSF

Roche Protease inhibitor tablets (containing EDTA)

BME, PMSF, and inhibitor tablets were added freshly.

Wash Buffer II:

15 50-200 mM HEPES pH 7.5 50-300 mM NaCl 1-15% Glycerol

Elution Buffer:

20 50-200 mM HEPES pH 7.5 50-200 mM NaCl 20-40% Glycerol

5-40 mM Glutathione (Reduced form)

Elution buffer should be about pH=7.5.

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6.3 Method of Making Proteins in a High-Throughput 96-Array Format

A yeast proteome microarray containing nearly all yeast proteins was prepared and screened for a number of biochemical activities. A high-quality collection of 5800 yeast ORFs (93.5% of the total) was cloned into a yeast high-copy expression vector using recombination cloning (Mitchell et al., 1993, Yeast 9:715). The yeast proteins were fused to GST-HisX6 at their amino termini and expressed in yeast under the control of a galactose-inducible *GAL1* promoter (Zhu et al., 2000, Nat. Genet. 26:283; Mitchell et al., 1993 Yeast 9:715). The yeast expression strains contain individual plasmids in which the correct yeast ORFs have been shown to be properly fused in-frame to GST by DNA sequencing. Briefly, yeast ORFs were amplified by PCR and co-transformed into yeast cells

along with the vector to generate expression clones. The plasmids were rescued in *E. coli*, and the vector-insert junctions were sequenced (FIG. 1A). If the ORF cloned was not the ORF of interest, or a frameshift was detected, the cloning cycle was repeated. Once a construct was confirmed, the plasmid DNA was reintroduced into yeast and *E. coli* to create permanent stocks for future analyses (Zhu et al., 2000, Nat Genet. 26:283). By repeating the cloning cycle, 5800 unique yeast ORFs were successfully cloned, representing 93.5% of the total.

To generate purified proteins for biochemical analysis, a robust and high-throughput purification method for preparing proteins in a 96-well format was developed and optimized. Using glutathione-agarose beads, yeast extracts were prepared, and fusion proteins were purified (for details of 96-well format protein purification protocol, a full list of results from all the experiments, and the design of the positive identification algorithms, visit public web site spine.mbb.yale.edu/protein_chips/). The lysis buffer and initial washes contained 0.1% Triton to ensure that the purified proteins were free of lipids. Using the procedures of the invention, at least 1152 protein samples can be prepared from cells in under 10 hours.

The quality and quantity of the purified proteins were monitored using immunoblot analysis of 60 random samples (FIG. 1B). Greater than 80% of the strains produced detectable amounts of fusion proteins of the expected molecular weight. A manual printing tool was used to spot 3 nl of 1152 purified proteins in duplicate onto glass slides (19), and the proteins detected using polyclonal anti-GST antibodies. Greater than 85% of the samples exhibited a visible signal above background, consistent with the immunoblot analysis. Using the procedures of the invention, fusion proteins from 6144 (64X96-well boxes) yeast strains can be purified in under two weeks.

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6.4 Method of Making a Proteome Microarray

To prepare the proteome chips, 6566 protein preparations representing 5800 different yeast proteins were printed in duplicate onto glass slides using a commercially available microarrayer. As a control, known amounts of GST were also printed. Two types of slides were employed. Aldehyde-treated microscope slides were used in initial experiments (MacBeath et al., 2000, Science 289:1760), in which case fusion proteins were attached to the slide surface through primary amines at the N-termini or other residues of the fusion proteins, resulting in a relatively random orientation of proteins on the surface. Proteins were spotted onto nickel-coated slides in subsequent experiments. In this case, fusion proteins are attached through their HisX6 tags such that the cloned yeast portions of

the fusion proteins are essentially uniformly oriented away from the surface. Although both types of slides were successfully used, the nickel-coated slides gave significantly superior signals for our particular protein preparations (FIG. 2A).

To determine how much fusion protein was covalently attached to different glass surfaces, and assess the reproducibility of the protein attachment, chips were probed with anti-GST antibodies. Over 93.5% of the protein samples gave signals significantly above background (i.e., greater than approximately 10 fg of protein). A comparison with known amounts of GST also printed on the slide, indicated that about 90% of the spots contain approximately 10 fg to 950 fg of protein. FIG. 2A shows that detection of proteins on a proteome chip with fluorescently labeled antibodies is extremely sensitive, i.e., the signal-to-noise ratio is high despite that only 1/10,000 of purified proteins from a 3-ml culture is spotted on the slide. The results demonstrate that it is feasible to spot 13,000 protein samples in one half the area of a standard microscope slide (2.5 cm by 7.5 cm) with excellent resolution (FIGS. 2A and 2B). To test the reproducibility of the protein spotting, 15 the signals from each pair of duplicated spots were compared with one another. As shown demonstrated by the sharp spike in FIG. 2C, 95% of the signals were within 5% of the average (10).

6.5 Method of Using a Proteome Microarray

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20 Proteome chips were tested by probing for several exemplary types of biological activities: protein-protein interactions, protein-nucleic acid interactions, and protein-lipid interactions.

Generally, proteome chips were prepared for assays as follows. The proteome chips were blocked by slowly immersing the printed glass slides into either BSA (1-3% (w/w) 25 BSA in PBS buffer; SIGMATM, USA) or glycine blocking buffer (30-300 mM glycine; 50-300 mM Tris, pH 6.5-8.5; 50-300 mM NaCl; SIGMATM, USA) with the protein side up. The buffer was filtered through a 2 µm filter unit to remove particles. Glycine is preferable when probing for carbohydrate-binding proteins. The slides were incubated in the blocking buffer at 4°C overnight without any shaking (disturbance of the blocking buffer may result in the protein streaks on the glass surface).

Probe proteins were generally prepared as follows. Yeast proteins were purified by affinity column using glutathione beads from 50 ml culture using standard protocols without the elution from the beads. The protein beads were washed three to five times with cold PBS buffer (pH 8.0) (SIGMATM, USA). Approximately 1 ml of Sulfo-NHS-LC-LC-Biotin 35 (PIERCETM Cat No. 21338, USA) dissolved in PBS (pH 8.0) at a concentration of 0.1-50

mg/ml was added to the glutathione beads and incubated at 4°C for 2h. The beads were washed 5 times with cold PBS buffer (pH 8.0) and eluted with 100-500 ml of the elution buffer (50-200 mM, HEPES pH 7.5; 50-200 mM NaCl; 20-40% glycerol; 5-40 mM glutathione). Protocols resulting in more weakly biotinylated proteins are preferred.

5 Batches of proteins that are biotinylated to different degrees were pooled for future usage.

6.5.1 Identification of Calmodulin-Interacting Proteins

To test for protein-protein interactions, the yeast proteome was probed with calmodulin (11). Calmodulin is a highly conserved calcium-binding protein involved in many calcium-regulated cellular processes and has many known physical partners (Hook et al., 2001, Annu. Rev. Pharmacol. Toxicol. 41:471). The calmodulin probe was biotinylated, and bound probe was detected using Cy3-labeled streptavidin. As a control, the yeast proteome was probed with Cy3-labeled streptavidin alone.

Generally, protein-protein interactions can be assayed as follows. Blocked proteome chips are washed three to five times in PBS buffer, and the extra liquid on the glass surface is removed by tapping the slides vertically on a KIMWIPETM. 200 μl of biotinylated protein probe is added to the proteome chip and immediately covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). After trapped air bubbles are removed, the chip is incubated in a humidity chamber at room temperature (RT) for one hour. To remove the cover slip, the chip is immersed into large volume of PBS buffer (>50 ml), and the slip should float away. The chip is then moved to a second PBS bath (>50 ml) and washed 3 X 5 min with shaking at RT. After removing extra liquid on the surface of the chip, at least 150 μl of Cy3- or Cy5- conjugated streptavidin (PIERCETM, USA; 1:2000 to 1:4000 dilution) is added to the surface and covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). The chip is incubated for greater than 30 min in the dark at RT. The chip is then washed as described above. To completely remove the liquid on the chip, the chip is spun to dryness at 1500-2000 rpm for 5-10 min at RT.

If a proteome chip is to be screened with antibodies, the protein-antibody interaction can be detected as follows. Blocked proteome chips are washed three to five times in PBS buffer, and the extra liquid on the glass surface is removed by tapping the slides vertically on a KIMWIPETM. 200 µl of primary antibodies (properly diluted in PBS containing 1-3% BSA and 0.1% TritonX-100) is added to the proteome chip and immediately covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). After trapped air bubbles are removed, the chip is incubated in a humidity chamber at RT for one hour. To remove the cover slip, the chip is immersed into large volume of PBS buffer (>50 ml), and the slip

should float away. The chip is then moved to a second PBS bath (>50 ml) and washed 3 X 5 min with shaking at RT. After removing extra liquid on the surface of the chip, >150 µl of Cy3- or Cy5-conjugated secondary antibodies (properly diluted in PBS containing 1-3% BSA and 0.1% TritonX-100) is added to the surface and covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). The chip is incubated for greater than 30 min in the dark at RT. The chip is then washed as described above. To completely remove the liquid on the chip, the chip is spun to dryness at 1500-2000 rpm for 5-10 min at RT.

When biotinylated calmodulin was used to probe the proteome chips in the presence of calcium, six known calmodulin targets, namely Cmk1p, Cmk2p, Cmp2p, Dst1p, Myo4p, 10 and Arc35p were identified (FIG. 3A). Cmk1p and Cmk2p are type I and II calcium/calmodulin-dependent serine/threonine protein kinases, which are both involved in the signal transduction pathway in the mating response (Hook et al., 2001, Annu. Rev. Pharmacol. Toxicol. 41). Cmp2p (Cna2p) is one of the two yeast calcineurins, and has been demonstrated to interact with calmodulin in vivo (Cyert et al., 1991, Proc. Natl. Acad. Sci. 15 U.S.A. 88:7376), Dst1p plays a role in transcription elongation (Stirling et al., 1994, EMBO J. 13:4329), Myo4p is a class V myosin heavy chain required for proper localization of ASH1 transcript (Bohl et al., 2000, EMBO J. 19:5514; Bertrand et al., 1998, Mol. Cell 2:437). Arc35p is a component of the Arp2/3 actin-organizing complex, which is involved in actin assembly and function, and in endocytosis (Winter et al., 1999, Proc. Nat. Acad. 20 Sci. U.S.A. 96:7288). Arc35p was recently shown to interact with calmodulin in a two-hybrid study (Schaerer-Brodbeck et al., 2000, Mol. Biol. Cell 11:1113), thus confirming the data herein demonstrating that Arc35p and calmodulin interact in vitro. Of the six known calmodulin targets that were not detected, two were not represented in the collection and the remaining four were not detectable in the GST probing experiments. In 25 addition to known interactors, the calmodulin probe identified 33 additional interactors. These interactors include a wide variety of different types of proteins (see Table 1; public web site bioinfo.mbb.yale.edu/proteinchip), consistent with a role for calmodulin in many

In addition to the calmodulin-binding targets, one protein, Pyc1p, which bound 30 Cy3-labeled streptavidin, was also identified. Pyc1p encodes a pyruvate carboxylase 1 homolog that contains a highly conserved biotin-attachment region (Menendez et al., 1998, Yeast 14:647). Thus, as predicted by its sequence, Pyc1p is biotinylated *in vivo*; and was identified in all experiments using streptavidin detection methods. Thus, proteome microarrays can be used to identify posttranslational modifications of proteins.

diverse cellular processes.

6.5.2 Identification of a Calmodulin-binding Motif

To identify putative calmodulin-binding domains, amino acid sequences shared among the different calmodulin-binding targets (*i.e.*, interactors) were determined (Zhu et al., 2000, Nat. Genet. 26:283). Fourteen of the 39 calmodulin-binding proteins contain a motif whose consensus is I/L-Q-X-X-K-K/X-G-B (SEQ ID NO: 1), where X is any residue and B is a basic residue (FIG. 3B). A related sequence in myosins, I-Q-X-X-X-X-X-X-X-R (SEQ ID NO: 16), has been shown previously to bind calmodulin (Homma et al., 2000, J. Biol. Chem 275). The results demonstrate that the motif, I-Q-X-X-X-X-X-X-X-X-R (SEQ ID NO: 16), is found in many calmodulin-binding proteins. Other calmodulin-binding interactors that lack this motif can possess other calmodulin-binding sequences.

6.5.3 Identification of a ATP/GTP-binding Proteins

- 1. Blocked proteome chips are washed three to five times in cold PBS buffer, and the extra liquid on the glass surface is removed by tapping the slides vertically on a KIMWIPETM.
 - 2. 100 μl of ATP and GTP solution (0.5-5 mM ATP-Cy3, 0.5-5mM GTP-Cy5, 100-400 mM NaCl, 1-30 mM MgCl₂, 50-300 mM Tris, pH 7-8.5) is added to the proteome chip and immediately covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, OR, USA). After trapped air bubbles are removed, the chip is incubated in a humidity chamber at 4°C for one hour.

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- 3. To remove the cover slip, the chip is immersed into large volume (>30 ml) of ice cold wash buffer (100-400 mM NaCl, 1-30 mM MgCl₂, 50-300 mM Tris, pH 7-8.5), and the cover slip should float away. The chip is then moved to a second cold wash bath (>50 ml) and washed 3 X 5 min with shaking at 4°C.
- 4. After removing extra liquid on the surface of the chip, the chip is spun to dryness at 1500-2000 rpm for 5-10 min at 4°C.

6.5.4 Identification of DNA-binding Proteins

Protein-DNA and protein-RNA interactions are important for many fundamental biological functions such as transcription regulation, chromosome segregation and maintenance, and RNA transport and processing (Williamson, 2000, Nat. Struct. Biol. 7(10):834-7). To explore the possibility of using a proteome chip to identify proteins that bind to DNA or RNA molecules, total yeast genomic DNA was labeled using Cy3-CTP and

Klenow fragment, and polyA⁺ was labeled using a biotinylated psoralen-derivative (31). Each probe was incubated with a different proteome chip (FIG. 3A).

The protein-nucleic acid interaction was assayed as follows. The nucleic acid probes were labeled as described by Winzeler et al. (1999, Science 285:901). Blocked proteome chips were washed three to five times in PBS buffer, and the extra liquid on the glass surface was removed by tapping the slides vertically on a KIMWIPETM. 200 µl of labeled nuclei acid probe (50-200 mM poly(dC/dG); 100-300 mM NaCl; 50-300 mM HEPES, pH 7.0-8.5; 10 mM MgCl₂) was added to the proteome chip and immediately covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). After trapped air bubbles were removed, the chip was incubated in a humidity chamber in the dark at room temperature ("RT") for one hour. To remove the cover slip, the chip was immersed into large volume of PBS buffer(>50 ml), so that the slip would float away. The chip was then moved to a second PBS bath (>50 ml) and washed 3 X 5 min with shaking in the dark at RT. To completely remove the liquid on the chip, the chip was spun to dryness at 15 1500-2000 rpm for 5-10 min at RT.

Visual inspection revealed that the genomic DNA probe identified 41 DNA-binding proteins (*see* Table 1). These included eight previously known DNA-associated proteins and two predicted DNA-binding proteins. Of the known DNA-binding proteins, we found three DNA-repair proteins (*i.e.*, xrs2p, Cdc1p, and Rad5lp (Costanzo et al., 2001, Nucleic Acids Res. 29:75)), a SWI-SNF global transcription activator complex component (*i.e.*, Snfl lp (Costanzo et al. 2001, Nucleic Acids Res. 29:75)), the alpha-subunit of the NC2 (Dr1/Drap1) repressor of class II transcription Bur6p (Sternberg. et al., 1987, Cell 48:567), the transcription factor Spt2p (a negative regulator of HO gene transcription (Sternberg et al., 1987, Cell 48:567)), and Met30p which contains five WD-repeats and binds certain promoter regions (Thomas et al., 1995, Mol. Cell Biol. 15:6526). We also identified Trm2p, which binds both DNA and RNA and is involved in DNA repair and RNA processing/modification (Sadekova et al., 1996, Curr. Genet. 30(1):50-5.).

Eight proteins, identified as nucleic-acid binding proteins, have unknown functions. Of these, three are likely to possess DNA-binding activities based on their amino acid sequences and the results of the instant assay. Ycr087c contains a zinc-finger domain, a common DNA-binding motif. Yhr054c shares sequence homology with the transcription factor Rsc3p (Costanzo et al., 2001, Nucleic Acids Res. 29:75). Ybr025c is believed to have putative nucleotide-binding activity (Costanzo et al., 2001, Nucleic Acids Res. 29:75). The instant assay shows that these eight proteins bind DNA *in vitro* and likely bind nucleic acids *in vivo*.

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FARLE 1

"-" indicates spots that are flagged as bad, which were given the value -10000 in the data table.

ORF	Block	Column	Row	· PC	PI(3,4,5)P ₃	PI(3,4)P ₂	PI(3)P	PI(4,5)P ₂	PI(4)P
MPS1	7	13	17	4374.59	1864.12	4216.40	310.00	1084.79	6148.60
YGR052W	13	13	17	2177.02	666.22	2328.28	00.6	563.52	164.18
YLL023C	47	. L'	18	1798.73	635.18	390.40	l	1868.44	1712.76
SNF1	41	15	17	1774.37	560.70	298.26	202.00	423.73	1795.06
YFL068W	11	6	4	1541.05	369.22	90.068	82.75	190.96	.931.28
GCNZ	25	15	17	1087.04	975.60	232.01	26.50	505.60	1356.97
YFR053C	7	13	4	1077.95	235.69	713.77	238.75	613.81	991.38
CMK1	13	15	17	1073.54	. 20.669	1786.71	7.00	545.23	105.39
YGL059W	21	15	17	970.24	273.38	3088.36	91.50	299.61	924.10
CAK1	41	13	17	854.02	269.10	245.20	99.50	233.85	1431.44
BUD32	45	13	17	621.00	108.67	249.28	318.00	282.63	666.29
SPS1	17.	13	17	557.02	389.70	1147.50	88.50	358.84	246.92
<u>TOS3</u>	37	15	17	535.89	623.47	766.36	167.00	685.45	2026.31
APG1	45	15	17.	464.28	92.70	138.45	72.00	319.65	668.47
LYS1	14	1	9	255.33	61.20	81.94	3.50	22.21	198.15
YHR174W	34	5	9	250.34	80.55	274.24	19.25	47.69	192.48
OPY1	23	15	1	250.04	41.17	179.27	17.50	37.89	188.13
RIM15	5	15	17	244.76	1219.28	4941.00	52.50	687.19	2349.00
YBR028C	3	13	17	232.43	147.60	225.42	18.00	165.05	209.03

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	PI(4)P	288.29	'	153.94	189.44	226.45	175.94	109.31	158.95	68.81	85.35	88.84	19.81	111.70	84.92	52.04	183.34	135.44	98.64	219.92	36.15	12.63	62.06
5	PI(4,5)P ₂	116.71	-4.57	26.56	57.48	18.73	244.09	101.47	69.63	45.07	112.57	123.90	ŧ	49.43	67.06	63.58	40.94	-	32.01	152.85	67.28	1	47.90
1(PI(3)P	51.50	17.00	35.25	19.50	20.00	117.00	28.00	•	17.75	42.50	-0.50	1	0.75	33.50	33.25	24.00		51.00	·	34.75	15.50	5.75
.15	PI(3,4)P ₂	192.45	230.60	199.20	132.49	ŧ	95.44	264.35	10.05	76.13	85.40	78.02	48.66	26.53	234.84	137.98	220.55	28.26	29.20	15.54	140.65	128.72	48.51
	PI(3,4,5)P ₃	115.88	166.05	14.06	93.38	78.07	25.42	94.05	56.47	87.75	66.04	67.95	86.09	33.86	65.47	99.11	97.20	37.12	30.15	21.38	48.26	37.12	25.31
20	PC	221.87	220.99	214.24	207.20	193.11	188.12	174.33	160.24	140.58	137.64	134.71	127.66	127.37	123.26	121.50	121.50	120.33	119.74	118.57	109:17	108.59	107.41
	Row	2	9	3	17	2	1	16	17	4.	10	10	6	4	17	9	3	3	3	18	3	1	4
25	Column	11	7	5	13	3	13	1	7	11	7	5	5	. 6	15	6	11	13	13 .	6	5	15	6
30	Block	32	17	24	11	18	47	4	30	2	44	44	76	25	7	2	11	38	34	47	2	1	21
35	ORF	YCL075W	<u>SLS1</u>	COX20	AKL1	HIS7	MET8	<u>YMR172C-A</u>	PAT1	YEL048C	GPM3	YOL117W	DFG5	YFR020W	KCC4	<u>RÓT1</u>	YDR288W	BTT1	ARP10	YLR022C	YDR070C	YBL107W-A	FBP26

35	30	25		20	15	15	10	5	
ORF	Block	Column	Row	PC	PI(3,4,5)P ₃	PI(3,4)P ₂	PI(3)P	PI(4,5)P ₂	PI(4)P
MY04	9	15	17	103.89	51.07	171.73	2.00	4.79	36.58
YER152C	22	5	4	103.60	50.17	24.80	-3.75	20.47	104.95
<u>FUR1</u>	4	11	9	95.67	72.22	210.19	112.25	111.70	148.94
YPL280W	28 ·	7	17	88.63	43.88	30.14	,	ı	118.45
YBR077C	17	13	1	86.58	•	129.98	4.75	21.12	66.85
TAT2	12	<i>L</i>	10	83.93	68.29	62.48	60.50	85.79	51.39
YIR041W	22	5	9	75.72	130.39	38.15	6.50	6.53	200.10
YKL031W	32	11	18	75.72	39.71	86.69	36.25	58.14	165.48
YGR242W	25	3	5	75.13	35.77	,11.93	17.00	10.02	38.32
HOR2	34	13	4	74.84	4.16	-3.30	11.00	7.19	68.15
<u>TAF25</u>	4	7	17	73.37	54.67	110.51	-10.00	77.08	45.29
<u>YJL193W</u>	33	13	9	72.78	•	21.03	97.00	27.65	103.43
MIS1	16	7	17	71.02	43.65	86.34	3.50	1	65.32
PEX4	16	5	5	69.85	21.49	8.63	1.75	28.31	57.48
SUN4	46	6	6	69.55	33.19	20.88	,	66.41	60.75
YOL073C	12	5	10	29.89	17.32	41.44	1.75	26.56	40.06
<u>A1</u>	4	3	16	68.38	48.49	88.38	31.75	46.81	31.57
YML058C-A	3	5	8	66.33	40.16	76.29	20.50	50.52	34.19
RPG1	17	6	Ţ.	66.33	22.05	61.53	-2.25	-6.75	54.00
VTC4	40	6	9	66.03	25.31	30.45	8.50	26.78	20.66
YDR149C	16	11	3	65.74	24.86	73.62	37.50	37.02	46.81
FRM2	2	15	17	63.39	16.88	59.34	3.50	75.34	12.63

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35	30	25		20		15	10	5	
ORF	Block	Column	Row	PC	PI(3,4,5)P ₃	$PI(3,4)P_2$	PI(3)P	PI(4,5)P ₂	PI(4)P
WRS1	76	5	10	62.51	51.08	63.10		21.56	4.57
<u>DPS1</u>	47	15	18 ·	62.22	13.05	25.12	ŧ	41.81	99.29
<u>PKR1</u>	47	5	8	61.63	15.75	37.52	t	87.75	61.40
YDR067C	2	6	3	61.34	-29.47	29.83	32.75	47.90	17.42
YCR025C	8	5	2	60.46	73.58	174.24	00.09	68.37	54.87
<u>APS2</u>	47	15	9	58.70	15.97	56.51	39.50	103.65	159.82
<u>RRP5</u>	48	5	16	58.11	21.26	22.60	ŧ	73.38	60.31
YMR148W	12	7	6	56.93	50.17	23.69	36.50	43.33	23.08
RPH1	14	13	2	56.93	9.90	90.73 · .	00.9	18.73	66.19
RGD1	36	15	2	54.59	19.35	54.31	00.9	55.31	127.60
YMR244W	44	5	6	54.29	28.24	29.20	64.75	44.20	32.66
SAS3	9	3	17	54.00	18.68	55.57	2.00	17.85	16.55
SGA1	16	3	16	52.83	26.55	60.59	16.50	6.75	41.15
SXMI	30	11	13	51.95	23.40	19.15		27.22	30.48
MUM2	3	13	1	46.66	63.34	49.29	81.50	38.76	90.15
VMA13	4	5	17	45.78	28.12	44.27	11.50	34.40	16.55
<u>YJL185C</u>	6	11	5	45.49	50.96	127,47	-3.25	13.94	27.00
YOL119C	44	1	10	44.61	14.62	5.97	360.50	23.08	22.65
SPS2	47	11	3	44.61	25.31	54.63	18.50	64.02	47.47
<u>YBL107C</u>	3	.15	1	44.02	65.93	23.86	104.50	34.84	40.50
<u>TH113</u>	2	11	3	43.14	39.04	75.66	13.75	64.67	55.09
KCS1	16	5	17	42.85	36.22	60.91	0.00	1	35.27
*									

5	$PI(4,5)P_2$ $PI(4)P$	43.98	50.30	35.71	87.31	32.88	- 16.11	43.98	8.06	31.35	101.90	28.74	38.76	42.24	32.88	37.45	-1.96	-4.79	-6.31	31.35	- 23.52	7 14.37	
		34.84	15.90	20.69	44.85	24.39		37.23	3.70	40.06	15.24	47.90	34.84	20.47	16.11	8.71	11.54	3.05	3.48	-1.31		237.77	
10	PI(3)P	22.50	3.50	0.25	-50.50	26.25	1.00	71.00	9.00	132.50	4.00	-15.50	ŧ	87.50	ī	•	51.25	23.00	213.00	-3.00	ı	1	
15	PI(3,4)P ₂	31.40	70.80	10.67	119.93	93.09	21.03	47.72	26.84	46.47	19.78	32.81	0.31	34.53	-2.83	,31.71	1	2.83	1.73	-5.02	7.53	68.6	
	PI(3,4,5)P ₃	40.05	33.41	11.02	13.05	90.90	8.55	27.45	26.89	39.15	21.15	19.12	16.20	29.25	-4.05	15.07	•	13.72	-6.08		33.97	2.59	
20	PC	42.26	41.97	39.62	39.33	38.15	38.15	37.86	31.70	31.10	31.11	29.05	28.76	28.17	26.12	25.24	24.36	24.07	23.77	22.30	22.30	21.72	
	Row	16	9	18	4	13	16	3	1	3	5	10	17	2	5	10	6	17.	10	4	9	10	
25	Column	15	3	7	3	13	15	13	5.	15	1	11	L	15	5		5	15	3	15	Ţ.	13	
30	Block	16	7	39	44	3	20	43	29	4	41	44	47	28	48	20	34	24	44	17	30	48	1
35	ORF	APA2	HYS2	DRS1	<u>GPA2</u>	SLT2	HEM14	TFB3	PCL6	YDL237W	YHL.041W	YOL054W	<u>YPL150W</u>	ARO4	<u>YGR176W</u>	MPD2	<u>GAD1</u>	DTR1.	ARG1	PAU5	CCT8	YPL275W	

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	PI(4)P	34.40	5.23	21.99	10.45	1	57.92	60.97	16.33	16.98	21.34	5.66	-1.52	29.18	-5.66	10.45	12.85	18.29	0.44	12.19	-3.92	25.26	-0.22
5	5)P ₂	. 3.	5.	2	-	•	5,)9	1	10	2	5.	-	25	3	12	12	18	0.	12	-3	25	위
	PI(4,5)P ₂	23.95	10.45	25.69	-2.61	26.13	25.69	89.69	18.73	1.31	5.23	14.81	16.33	5.66	15.68	-13.06	1.96	59.23	14.37	12.41	6.53	22.21	•
10	PI(3)P	11.50	21.50	f	-11.50	¢	43.50	. €	8.50	ſ	6.50	49.00	99.00	•	3.75	-9.00	1	-22.50	-6.50		-5.00	ı	1
15	PI(3,4)P ₂	-2.83	27.00	-0.16	87.91	-5.81	53.69	19.78	16.80	6.28	-2.20	1.57	7.53	5.34	2.67	-5.02.	7.38	-0.63	18.99	0.16	-3.14	6.59	-0.78
	PI(3,4,5)P ₃	42.86	14.62	4.16	0.23	17.66	27.00	12.60	16.65	4.27	29.02	-13.05	29.92	7.65	1.24	-1.12	-0.45	0.45	-9.22	-5.29	-0.90	8.77	3.38
20	PC ·	20.25	18.78	18.49	16.43	14.97	14.67	14.09	13.50	12.91	10.86	10.57	10.27	7.63	7.34	6.46	6.46	5.87	5.58	5.58	5.28.	5.28	4.99
	Row	9	11	10	17	1	3	3	5	16	4	13	13	10	7	1	6.	9	6	1	4	17	6
25	Column	13	15 ·	13	13	3	15	1	7	15	6	15	13 .	15	13	15	5	15	5	5	.15	7	9
30	Block	29	28	47	5	30	41	48	12	39	14	37	37	42	43	34	48	34	9	48	12	48	42
35	ORF	URA8	YPL107W	CAM1	KSS1	PET9	YDR348C	KGD2	YGL165C	SSB1	MAGI	ASF1	YKL121W	YOL053W	YLR020C	PYK1	RPL9B	APG7	SP01	RTG3	YEL064C	BPL1	GFD1

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	PI(4)P	32.66	-6.97	23.52	34.40	10.89	19.38	28.09	2.61	28.74	3.05	13.28	19.16	56.61	-9.15	32.66	70.89	-3.27	41.81	28.74	11.76	36.58
5	$PI(4,5)P_2$	13.50	-11.76	21.34	35.06	-8.27	-4.79	46.38	-8.06	21.99	-7.62	14.15	6.31	76.78	8.93	3.27	10.89	18.94	16.11	9.36	4.35	34.40
10	PI(3)P	-1.50	4.50	54.50	ı	51.50	-6.00	•	-47.75	28.75	ı	2.00	1	42.00	0.50	5.25	12.00	15.75	-5.00	-13.75	ı	•
	PI(3,4)P ₂	78.17	1	18.84	21.19	8.79	68.6	11.93	-8.95	19.62	-8.63	4.71	11.62	65.93	-11.93	106.12	35.48		15.85	4.55	3.45	15.54
15	PI(3,4,5)P ₃	32.17	-4.05	13.95	25.31	22.05	9.79	26.44	-8.66	13.16	-6.53	-2.14	5.62	55.35	-39.15	71.10	14.40	1.35	34.31	18.79	29.67	14.29
20	PC	3.52	3.52	3.52	3.23	2.93	2.64	0.29	-0.88	-1.17	-1.76	-4.70	-18.20	1	1	1	•	1	ŧ	1	ı	
	Row	9	5	3	6	2	8		7	6	18	18	4	2	1	9	3.	5	6	6	4	9
25	Column	1	15	15	9	13	6	13	11	5	6	11	. 6	11	3	5	15	9	3	5	15	13
	Block	7	40	43	40	28	29	48	44	40	38	43	18	2	17	17	19	34	37	37 ·	42	42
35	ORF	YJR080C	YGR101W	MRP1	YNL051W	STP22	<u>MUB1</u>	<u>ATP1</u>	CSE4	VAC7	GFA1	SDS22	YER046W		YBR025C	RFA3	MHR1	ERV1	<u>TOS6</u>	YNR069C	SDS3	YIL003W
	<u></u>	[>]	M	2	>	Ø		A	<u>ات</u>	>		59 -	Ι',		7	2	$ \Sigma $	国	Ĕ	>	S	

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6.5.5 Identification of RNA-binding Proteins

In addition to Trm2p, other interactors with RNA-binding activity were identified. For example, an isoleucyl-tRNA synthetase (IIslp) and a ribosomal protein (Rp141B) have both been shown to bind RNA molecules (Lanker et al., 1992, Cell 70:647; Suzuki et al., 1990, Curr. Genet 17:185). Other proteins involved in translation tested positive in the screen for RNA-binding proteins such as Rp141b, Sqt1p, and Gcd1 lp. Rpl41b is a component of the large ribosomal subunit. Sqt1p interacts with Rpll0p, which is an RNA-binding protein (Costanzo, 2001, Nucleic Acids Res. 29:75). Gcd1 lp, a gamma subunit of translation initiation factor eIF2 has GTP-binding activity, and binds to the Lsm1-7 complex (U6-specific snRNP) as demonstrated by two-hybrid analysis (FromontRacine et al., 2000, Yeast 17:95). Two other protein targets that bind the DNA

Lsm1-7 complex (U6-specific snRNP) as demonstrated by two-hybrid analysis (FromontRacine et al., 2000, Yeast 17:95). Two other protein targets that bind the DNA probe also bind nucleotides, *i.e.*, Thi13p is involved in nucleotide metabolism, and Ypt31p has GTP-binding activity (Costanzo et al., 2001, Nucleic Acids Res. 29:75). The RNA-binding and nucleotide-binding proteins identified with the DNA probe may reflect

15 low affinity interactions with DNA or, alternatively, may represent normal affinity for DNA that has been previously unrecognized.

RNA-polyA⁺ from exponentially growing cells was labeled with biotin and used to probe the proteome microarray. The positive signals were detected using Cy3-streptavidin. Nineteen target interactors were identified (*see* Table 1), including a U1 snRNP component (Smd1p), a protein known to bind mRNA import protein (Sxm8p), a C2-H2 zinc-finger protein (Azflp), and Tos8p, which was suggested by its sequence to be involved in RNA splicing and processing (Schwikowski et al., 2000, Nat. Biot. 18:1257). In addition to proteins that likely directly or indirectly interact with RNA, we found two transcription factors (Bur6p and Tos8p), one transcriptional silencer (Hst3p), six other proteins encoded by known ORFs, and seven proteins from uncharacterized ORFs. One of the unknown ORFs exhibited the highest signals (Yer152c). It is important to note that only one interactor, Bur6p, bound both the DNA and RNA probes. The other interactors were uniquely detected with either a double-stranded DNA probe or a RNA probe, suggesting that most proteins are likely to specifically bind either DNA or RNA.

In summary, of the 41 target interactors, eighteen are nucleic-acid interacting proteins; eleven are known or putative DNA-binding proteins, three are RNA-binding proteins, and four bind nucleotides.

6.5.6 Identification of Lipid-binding Proteins

The proteome chips of the invention are valuable for identifying activities that might not be accessible by other experimental approaches such as, for example, protein-drug interactions and protein-lipid interactions. Indeed, genome-wide analysis of proteins that 5 bind to phospholipids has not been explored previously for any species. A proteome chip of the invention was used to study proteins that interact with phosphatidylinositol ("PI"). In addition to their roles as constituents of cellular membranes, phosphatidylinositols are important second-messengers that regulate diverse cellular processes, including growth, differentiation, cytoskeletal rearrangements, and membrane trafficking, and are found in the 10 nucleus, vacuole, and plasma membrane (Odorizzi et al., 2000, TIBS 25:229; Fruman et al., 1998, Annu. Rev. Biochem. 67:481; Martin, 2000, Annu. Rev. Cell Dev. Bio. 14:231; Wera et al., 2001, FEMS Yeast Res. 1406:17). Because they are often present only transiently and in low abundance within cells, phosphatidylinositols have not been characterized extensively in yeast, and consequently, little is known about proteins that bind particular 15 phospholipids (Odorizzi et al., 2000, TIBS 25:229; Fruman et al., 1998, Annu. Rev. Biochem. 67:229; Martin, 2000, Annu. Rev. Cell Dev. Bio. 14:231; Wera et al., 2001, FEMS Yeast Res. 1406:1).

To identify PI-binding proteins in yeast, liposomes were used as probes because the liposome provides the most relevant physiological condition to assay the interactors. Six types of liposomes were used. Each contains phosphatidylcholine ("PC") with 1% (w/w) N-(biotinoyl)-1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (biotin DHPE). The biotinylated lipid serves as a label that can be detected by Cy3-streptavidin (21). In addition to PC, the five other liposomes contain either 5% (w/w) PI(3)P, PI(4)P, PI(3,4)P₂, PI(4,5)P₂, or PI(3,4,5)P₃ (FIG. 3A). Each of these phospholipids has been found in yeast except PI(3,4,5)P3 (Odorizzi et al., 2000, TIBS 25:229; Fruman et al., 1998, Annu. Rev. Biochem. 67:481; Martin, 2000, Annu. Rev. Cell Dev. Bio. 14:231; Wera et al., 2001, FEMS Yeast Res. 1406:1).

The protein-liposome interaction was assayed as follows. Appropriate amounts of each lipid in chloroform were mixed and dried under nitrogen. The lipid mixture was resuspended in TBS buffer by vortexing. The liposomes were created by sonication. Blocked proteome chips were washed three to five times in PBS buffer, and the extra liquid on the glass surface was removed by tapping the slides vertically on a KIMWIPETM. 200 µl of liposome solution was added to the proteome chip and immediately covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). After trapped air bubbles were removed, the chip was incubated in a humidity chamber at RT for one hour. To

remove the cover slip, the chip was immersed into large volume of PBS buffer (50 ml), so that the slip would float away. The chip was then moved to a second PBS bath (>50 ml) and washed 3 X 5 min with shaking at RT. After removing extra liquid on the surface of the chip, about 100 µl of Cy3 or Cy5 conjugated streptavidin (PIERCETM, USA; 1:2000 to 1:4000 dilution) was added to the surface and covered by a hydrophobic plastic cover slip (GRACE BIO-LABSTM, USA). The chip was incubated for greater than 30 min in the dark at RT. The chip was then washed as described above. To completely remove the liquid on the chip, the chip was spun to dryness at 1500-2000 rpm for 5-10 min at RT.

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The six liposomes identified a total of 150 different interactors that produced signals significantly higher than the background. An algorithm was devised to assist in the identification of positive signals (42). Fifty-two (35%) of the interactors (i.e., lipid-binding proteins) correspond to uncharacterized proteins thus ascribing the first biochemical activities to those proteins. These results also indicate that many previously uncharacterized proteins have potentially important biochemical activities.

Of the 52 uncharacterized proteins, thirteen (25%) are predicted to be associated with membranes (Gerstein, 1998, Proteins 33:518). Many others contain basic stretches, which can mediate electrostatic interactions with the negatively charged phosphatidylinositols.

Of the 98 previously described interactors, the subcellular localization of 81 of these
has been investigated (Costanzo et al., 2001, Nucleic Acids Res. 29:75). Most are either
membrane-associated proteins or are involved in lipid metabolism. More specifically, 45
proteins are membrane-associated and either have known or predicted membrane-spanning
regions (Gerstein, 1998, Proteins 33:518). Among these identified interactors are integral
membrane proteins, proteins with lipid modifications (e.g., the glycosylphosphatidylinositol
(GPI) anchor proteins Tos6p and Sps2p (Costanzo et al., 2001, Nucleic Acids Res. 29:75),
the prenylated proteins, Gpa2p and mating pheromone a-factor (Ansari et al., 1999, J. Bio.
Chem. 274:30052), and peripherally-associated proteins (e.g., Kcc4p and Myo4p which are
found at the bud neck and/or cell periphery (Bohl, 2000, EMBO J. 19:5514; Bertrand et al.,
1998, Mol. Cell 2:437)). Eight others are involved in lipid metabolism (e.g., Bpl1p),

inositol ring phosphorylation (e.g., Kcs1p), or predicted to be involved in membrane/lipid function (e.g., Ylr020cp which has homology to triacylglycerol lipase).

The interactors (*i.e.*, phospholipid-binding proteins) were grouped in several ways (42). First, the interactors were sorted according to whether they bound lipids strongly or weakly, based on the phospholipid-binding signal relative to the amount of GST (FIG. 4).

35 Slightly more (72%) of the strong lipid binding proteins (FIGS. 4A and 4B) were

characterized relative to the weakly binding proteins (54%) (FIGS. 4C and 4D). Many of the strong lipid-binding proteins are membrane-associated, or are predicted to be associated with membranes, whereas fewer of the weaker binding proteins appear to be associated with membranes (FIG. 4, "Membrane" column). Surprisingly, nineteen of the lipid-binding proteins are kinases, and seventeen of these are protein kinases. Moreover, thirteen of the seventeen protein kinases bind very strongly to the phospholipids.

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The proteins were also grouped by whether they preferentially bound one or more phosphatidylinositols as compared with phosphatidylcholine. One-hundred-and-one proteins bound to phosphatidylcholine as well or nearly as well as to the phosphatidylinositol lipids 10 tested (PI/PC < 1.3) (FIGS. 4B and 4D). However, 49 proteins bound to one or more phosphatidylinositols preferentially (PI/PC > 1.3) (FIGS. 4A and 4C). Analysis of the strong interactors (i.e., phosphatidylinositol-binding proteins) revealed that many specifically bound particular phosphatidylinositol lipids. For example, Stp22p, which is required for vacuolar targeting of temperature-sensitive plasma membrane proteins, such as 15 Ste2p and Can1p, preferentially binds PI(4)P (Li et al., 1999, Mol. Cell Biol. 19:3588). Nine protein kinases specifically bind PI(4)P and PI(3,4)P2 strongly and one binds these lipids weakly. Atp1p, the alpha subunit of F1-ATP synthase of the inner membrane of mitochondria, also preferentially binds PI(3,4)P2 (Arnold et al., 1999, J Biol. Chem). Sps2p, which is involved in middle/late stage of sporulation and is localized to the prospore 20 membrane (Chu et al., 1998, Science 282: 699), preferentially interacts with PI(3)P. One interesting example is Myo4p which preferentially binds PI(4,5)P2; perhaps binding of this lipid is an important part of its interaction at the cell cortex and/or its regulation. No strong lipid binding targets were found that specifically bound PI(3,4,5)P3 although some proteins bound both this lipid and others (FIG. 4). These results demonstrate that many 25 membrane-associated proteins including integral membrane proteins and peripherally associated proteins preferentially bind specific phospholipids in vivo.

Unexpectedly, many proteins involved in glucose metabolism, including regulators of glucose metabolism bind phospholipids. Specifically, three glucose metabolism enzymes, *i.e.*, phosphoglycerate mutase (Gpm3p), enolase (Eno2p) and pyruvate kinase (Cdc19p/Pyk1p) which participate in sequential steps in glycolysis, were identified. Hexokinase (Hxk1p), which converts glucose to glucose-6-phosphate, and two protein kinases known to be regulators of glucose metabolism (*i.e.*, Snf1p and Rim15p) were also identified as lipid-binding interactors. Hxk1p has recently been shown to bind zwitterrion micelles which stimulate its activity (30); Eno2p has been shown to be secreted suggesting that it might also interact with membranes (31). Accordingly, in one embodiment,

phospholipids can be used to regulate steps involved in glucose metabolism. In another embodiment, glycolytic steps can be enhanced by conducting steps of glucose metabolism on phospholipid scaffolds such as, but not limited to, cell membranes.

Surprisingly, the probes comprising phospholipids recognized many proteins not expected to be involved in membrane function or lipid signaling. Therefore, six proteins, i.e., Rim15p, Eno2p, Hxk1p, Sps1p, Ygl059wp, and Gcn2p, were further tested for phosphoinositol binding using two types of standard assays (Williamson, 2000, Nat. Struct Biol. 7(10):834-7). For Rim15p, Eno2p, and Hxk1p, PI(4,5)P2 liposomes were first adhered to a nitrocellulose membrane, which was then blocked by BSA; different amounts 10 of the GST fusion proteins and a GST control were used to probe the membrane, and bound proteins were detected using anti-GST antibodies. As shown in FIG. 5A, each yeast fusion protein tightly bound PI(4,5)P2 and exhibited a dosage effect; GST alone did not. We also carried out the reverse assay for four GST fusion proteins, Rim15p, Sps1p, Ygl059wp, and Gcn2p (Guerra et al., 2000, Biosci. Rep. 20: 41). Different amounts of these purified 15 proteins were spotted onto nitrocellulose filters and probed with the six different liposomes (FIG. 5B). The bound liposomes were detected using a horseradish peroxidase ("HRP")-conjugated streptavidin. As with experiments using the proteome chips, liposomes bound to each protein, and did not bind the BSA control. Sps1p bound all five phosphoinositol- containing liposomes nearly equally. Rim15p, Gcn2p, and Ygl059wp 20 exhibited different affinities to different liposomes (see FIG. 5B for Rim15p). All three proteins bound strongest to PI(3)P and PI(4)P and PI(3,4)P2. For Rim15p, a linear correlation between the binding signal and the level of Rim15p was revealed (FIG. 5C).

Further, several proteins with unknown functions also showed strong lipid-binding activities, indicating that they are likely to bind phospholipids in cells. Routine assays 25 known to those skilled in the art can be used to quickly screen lipid-binding interactors that are identified with a proteome chip to determine lipid-binding activity in vivo. Additionally, in accordance with the invention, a proteome chip can be used to determine if these interactors are involved in phospholipid metabolic pathways or signal transduction pathways.

In summary, these results unequivocally demonstrate that many lipid-binding proteins, including proteins not previously known to bind lipids, can be detected and characterized using a proteome chip of the invention. Moreover, phospholipid-binding interactors identified using the proteome chips of the invention can be shown to be bona fide lipid-binding proteins in conventional assays.

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Moreover, because such interactions cannot be studied by using either the yeast two-hybrid system or DNA/oligo microarray technology, the proteome chips of the invention and the methods of using them represent a pioneering approach to exploring molecule-protein interactions.

These studies indicate that it is feasible to prepare and screen a proteome microarray for a eucaryote. The novel and unobvious combination of several technological features were critical for practicing the present invention. First, the proteins analyzed in this study were prepared from yeast expression clones that have been verified by DNA sequencing and contain pure plasmids. Second, it was essential to produce defined clones and proteins in a 10 high throughput fashion. The procedures described herein provide enough protein for 5,000 protein chips. Third, the proteins were produced in a eucaryotic host. As such, large numbers of full-length proteins that are properly folded, posttranslationally modified and/or complexed with their native partners can be produced. As demonstrated by immunoblot analysis, at least 80% of the purified fusion proteins are expressed as full-length proteins.

Many interactors can bind to probes through associated proteins, and not directly. In some cases, the interactor can be part of a multimer. However, such associated proteins can also be detected and identified using the methods of the present invention. Nevertheless, the interactors detected by the methods of the invention likely directly interact with, or at least are tightly associated with a protein interacting with, a probe. Proteins of the proteome 20 chips of the invention were prepared using stringent conditions (e.g., washed with 0.5 M NaCl), and Coomasie staining revealed only those bands detectable by anti-GST antibodies indicating that contamination with other proteins was minimal.

The collection of proteins assembled is likely to underrepresent secreted proteins with properly folded extracellular domains. A GST tag and a HisX6 tag was fused 25 upstream of the translational initiation codon, such that membranous proteins having a signal peptide may not be delivered to the secretory pathway and may not be folded or modified appropriately. Three proteins having signal peptides were identified in the screens for lipid-binding proteins, suggesting that at least some secreted proteins are produced and contain functional domains.

30 Further, because not all proteins are readily overproduced and purified using the high-throughput methods practiced, not all interactions were detected. The protein arrays used are estimated to contain approximately 80% of the full-length yeast proteins at reasonable levels for screening, however, and thus most protein interactions can be expected to be detected.

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These results demonstrate that a proteome chip can be screened for biochemical activities, thereby allowing global proteome analysis. Similar procedures can be used to prepare protein arrays of 10-100,000 proteins for global high-throughput proteome analysis in humans and other eukaryotes.

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- GST gene and the polycloning site of pEG(KG). The yeast ORFs were cloned using the strategy described previously (5), except every step was done in a 96-well format. Plasmid DNAs confirmed by DNA sequencing were reintroduced into both yeast (Y258) and *E. coli* (DH5a). The library contains 5800 unique ORFs.
 - Details of a 96-well format protein purification protocol, a full list of results from all
- 25 the experiments, and the design of the positive identification algorithms, can be found at public web site bioinfo.mbb.yale.edu/proteinchip.
 - Biotinylated calmodulin (CalBiochem, USA) was added to the proteome chip at 0.02 mg/ml in PBS buffer with 0.1 mM calcium and incubated in a humidity chamber for one hour at room temperature. 0.1 mM calcium was present in buffers in all subsequent
- 30 steps. The chip was washed three times with PBS at RT. Cy3-conjugated streptavidin (Jackson IR, USA) (1:5000 dilution) was added to the chip and incubated for 30 min at RT. After extensive washing, the chip was spun dry and scanned using a microarray scanner; the data was subsequently acquired with the GENEPIXTM array densitometry software (Axon, USA).
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 - Liposomes were prepared using standard methods (52). Briefly, appropriate amounts of each lipid in chloroform were mixed and dried under nitrogen. The lipid
- 15 mixture was resuspended in TBS buffer by vortexing. The liposomes were created by sonication. To probe the proteome chips, 60 ml of the different liposomes were added onto different chips. The chips were incubated in a humidity chamber for one hour at RT. After washing with TBS buffer for three times, Cy3-conjugated streptavidin (1:5000 dilution) was added to the chip and incubated for 30 min at RT.
- 20 22 Positives were identified using a combination of the GenePix software which computes a local intensity background for each spot and a series of algorithms we developed. Details can be found at: public web site bioinfo.mbb.yale.edu/proteinchip.
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7. REFERENCES CITED

All references cited herein are incorporated herein by reference in their entirety and for all purposes to the same extent as if each individual publication or patent or patent application was specifically and individually indicated to be incorporated by reference in its entirety for all purposes.

8. EQUIVALENTS

Many modifications and variations of this invention can be made without departing from its spirit and scope. A person of ordinary skill in the art will recognize, or be able to ascertain through routine experimentation, various alternatives, adaptations, and modifications to the particular embodiments of the invention described herein, all of which are within the scope of the invention. Accordingly, the claimed invention intends to encompass all such equivalents. Thus, the specific embodiments described herein are offered by way of example only, and the invention is to be limited only by the terms of the appended claims, along with the full scope of equivalents to which such claims are entitled.

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We claim:

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1. A positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least one protein encoded by at least 50% of the known genes in a single species.

- 2. The array of Claim 1, wherein the plurality of proteins comprises at least one protein encoded by at least 70% of the known genes in a single species.
- 3. A positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 50% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein.
 - 4. A positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1000 proteins expressed in a single species.
- 5. A positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins in aggregate comprise proteins encoded by at least 1000 different known genes in a single species.
- 6. The array of Claim 1, 3, 4, or 5, wherein the proteins are organized on the array according to a classification of proteins.
- 7. The array of Claim 6, wherein the classification is by abundance, function, enzymatic activity, homology, protein family, association with a particular metabolic pathway, or posttranslational modification.
 - 8. The array of Claim 1, 3, 4, or 5, wherein the proteins are attached to the solid support via a His tag.

9. The array of Claim 1, 3, 4, or 5, wherein the solid support comprises nickel.

10. The array of Claim 1, 3, 4, or 5, wherein the solid support comprises a nickel-coated slide.

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11. A method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least one protein encoded by at least 50% of the known genes in a single species.

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12. A method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 50% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein.

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13. A method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins comprises at least 1000 proteins expressed in a single species.

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14. A method for making a positionally addressable array comprising the step of attaching a plurality of proteins to a surface of a solid support, with each protein being at a different position on the solid support, wherein the plurality of proteins in aggregate comprise proteins encoded by at least 1000 different known genes in a single species.

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15. A method for making a positionally addressable array comprising the step of attaching a plurality of fusion proteins to a surface of a solid support, with each fusion protein being at a different position on the solid support, wherein the fusion protein comprises a first tag, a second tag, and a protein sequence encoded by genomic nucleic acid of an organism.

5	16.	The method of Claim 15, wherein prior to said attaching step is a step of purifying the protein by contacting the protein with the binding partner of the said first tag, and wherein the second tag is used in said attaching step to attach the protein to the solid support.
3		attach the protein to the solid support.
	17.	The method of Claim 16, wherein the first tag is a GST tag and the second tag is a His tag.
10	18.	The method of Claim 15, wherein the first tag and the second tag are found at the amino-terminal end of the protein.
	19.	The method of Claim 15, wherein the first tag and the second tag are found at the carboxy-terminal end of the protein.
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	20.	A method for making and isolating a plurality of purified protein samples, comprising the steps of:
		at each site of a plurality of sites of a multi-site array:
		(a) growing a eukaryotic cell having a heterologous nucleotide sequence
20		operatively linked to a regulatory sequence;
		(b) contacting the regulatory sequence with an inducer that enhances
		expression of a protein encoded by the heterologous nucleotide sequence; (c) lysing the cell to produce a cell lysate;
		(d) contacting the cell lysate or protein-containing sample therefrom with a
25		binding agent such that a complex between said protein and binding agent is formed; and
		(e) isolating the protein from the complex;
		wherein each step is conducted in a multi-array format.
30	21.	The method of Claim 20, wherein each site is a well.
	22.	The method of Claim 20, wherein said protein is a fusion protein comprising an affinity tag to which said binding agent binds.
35	23.	The method of Claim 20, wherein the cell is a yeast cell.

24. The method of Claim 20, wherein said lysing step is performed using a paint shaker.

- A method for detecting a lipid-binding protein comprising the steps of:

 (a) contacting a probe comprising a lipid with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support; and

 (b) detecting any protein-probe interaction, wherein detection of the interaction at a position on the solid support indicates the presence of a lipid-binding protein at said position.
 - 26. The method of Claim 25, wherein the lipid is a phospholipid.
- 15 27. The method of Claim 26, wherein the phospholipid is phosphatidylcholine or phosphatidylinositol.
 - 28. The method of Claim 25, wherein the probe comprises a liposome.
- 29. A method for detecting a binding protein comprising the steps of:

 (a) contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least one protein encoded by at least 50% of the known genes in a single species; and

 (b) detecting any protein-probe interaction.
- 30. A method for detecting a binding protein comprising the steps of:

 (a) contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 50% of all proteins expressed in a single species, wherein protein isoforms and splice variants are counted as a single protein; and

 (b) detecting any protein-probe interaction.
- 35 31. A method for detecting a binding protein comprising the steps of:

(a) contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprises at least 1000 proteins expressed in a single species; and

- 5 (b) detecting any protein-probe interaction.
 - 32. A method for detecting a binding protein comprising the steps of:

 (a) contacting a probe with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins in aggregate comprise proteins encoded by at least 1000 different known genes in a single species; and (b) detecting any protein-probe interaction.
 - 33. A method for detecting a binding protein comprising the steps of:

 (a) contacting a probe with a positionally addressable array comprising a plurality of fusion proteins, with each fusion protein being at a different position on a solid support, wherein the fusion protein comprises a first tag, a second tag, and a protein sequence encoded by genomic nucleic acid of an organism; and
- 20 (b) detecting any protein-probe interaction.
 - 34. The method of Claim 29, 30, 31, 32, or 33, wherein the probe comprises a nucleic acid, protein, small molecule, drug candidate or lipid.
- 25 35. The method of Claim 34, wherein the nucleic acid comprises RNA or DNA.
 - 36. The method of Claim 34, wherein the probe is a yeast protein.
- 37. The method of Claim 36, wherein the yeast protein is Myo2, Rho1, Rho2, Rho3, Rho4, Cdc11, Cdc12, or Hs17.
 - 38. The method of Claim 34, wherein the probe is an antibody.

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39.	The method of Claim 38, wherein the antibody is directed against cyclin,
	kinase, GST, Clb5, Cla4, Ste20, Cdc42, PI(3,4)P2, PI(4)P, SPA2, CLB1,
	CLB2, or Cdc11.

- 5 40. The method of Claim 34, wherein the probe is calmodulin.
 - 41. The method of Claim 34, wherein the probe comprises a small molecule selected from the group consisting of ATP, GTP, cAMP, phosphotyrosine, phosphoserine, and phosphothreonine.
 - 42. The method of Claim 34, wherein the probe comprises phosphatidylcholine or phosphatidylinositol.
 - 43. The method of Claim 34, wherein the probe comprises a liposome.
 - 44. The method of Claim 25, 29, 30, 31, 32, or 33, wherein the probe is from a mammal.
 - 45. The method of Claim 44, wherein the mammal is human.
 - 46. The method of Claim 44, wherein the plurality of proteins is non-human.
 - 47. The method of Claim 25, 29, 30, 31, 32, or 33, wherein the plurality of proteins is attached to the solid support via a His tag.
 - 48. The method of Claim 25, 29, 30, 31, 32, or 33, wherein the solid support comprises nickel.
 - 49. The method of Claim 25, 29, 30, 31, 32, or 33, wherein the solid support comprises a nickel-coated slide.
 - 50. The method of Claim 34, further comprising the step of determining the identity of a probe whose interaction with a protein is detected in said detecting step.

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51. The method of Claim 50, wherein said interaction indicates that said identified probe is an antibacterial, antifungal, or antiviral protein.

- 52. A method of labeling a protein for use in a binding assay, comprising the steps of:
 - (a) contacting separate aliquots of said protein with a biotin-transferring compound under conditions and for a period of time to produce said proteins that are biotinylated to differing degrees among the different aliquots; and (b) combining together said different aliquots to produce a sample of differentially biotinylated protein.
 - 53. A method for detecting a binding protein comprising the steps of:

 (a) contacting a sample of biotinylated protein produced by the method of
 Claim 52 with a positionally addressable array comprising a plurality of
 proteins, with each protein being at a different position on a solid support;
 and
 - (b) detecting any positions on the array, wherein interaction between a biotinylated protein and a protein on the array occurs.
- 54. A method for detecting a binding protein comprising the steps of:

 (a) contacting a sample of biotinylated protein produced by the method of Claim 52 with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support;

 (b) contacting said array with streptavidin conjugated to a fluor; and

 (c) detecting any positions on the array at which fluorescence occurs, wherein said fluorescence indicates that interaction between a biotinylated protein and a protein on the array occurs.
 - A method for determining whether a protein preferentially binds phosphatidylinositol as compared with phosphatidylcholine, comprising the steps of:
 - (a) contacting a probe comprising phosphatidylinositol with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support;

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(b) detecting protein-probe interaction, wherein said interaction at a position on the solid support indicates the presence of a phosphatidylinositol-binding protein;

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(c) contacting a probe comprising phosphatidylcholine with a positionally addressable array comprising a plurality of proteins, said proteins comprising at least some of the same proteins as in step (a), with each protein being at a different position on a solid support;

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(d) detecting protein-probe interaction, wherein the interaction at a position on the solid support indicates the presence of a phosphatidylcholine-binding protein; and

(e) comparing, for each of a plurality of proteins, the results of steps (b) and (d).

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A method for determining if a phospholipid regulates a metabolic pathway or signal transduction pathway in a cell, or if said metabolic or signal transduction pathway occurs on membrane surfaces, comprising the steps of:

(a) contacting a probe comprising phospholipid with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support, wherein the plurality of proteins comprise one or more proteins that form at least part of said pathway; and (b) detecting interaction of said probe with a protein in said pathway; wherein said interaction indicates that said probe regulates said metabolic pathway or signal transduction pathway, or that said pathway occurs on membrane surfaces.

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57. A method for making a non-naturally occurring protein that binds calmodulin comprising making a non-naturally occurring protein comprising the following sequence:

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I/L-Q-X-X-K-K/X-G-B (SEQ ID NO: 1),

wherein X is any amino acid and B is a basic amino acid.

58. A method for determining the presence or absence of a posttranslational modification in a protein comprising the steps of:

(a) contacting a probe that binds to said posttranslational modification with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support; and (b) detecting any interaction of said probe with a protein; wherein said 5 interaction at a position on the solid support indicates that the protein at said position has said posttranslational modification. The method of Claim 58, wherein said posttranslational modification is 59. methylation, phosphorylation, biotinylation, acetylation, pegylation, 10 glycosylation, lipid modification, ubiquitination, and sumolation. A method for preparing a culture of yeast cells, comprising the steps of: 60. (a) growing a plurality of yeast cells in a growth medium until the OD_{600} is between 0.3 and 1.0, wherein said plurality of yeast cells comprises a 15 heterologous nucleotide sequence operatively linked to a regulatory sequence, (b) contacting said cell with an inducer that enhances expression of a protein encoded by said heterologous nucleotide sequence; (c) separating said cells from said medium; 20 (d) contacting said cells with cold water; (e) separating said cells from said cold water; (f) contacting said cells with cold lysis buffer; (g) separating said cells from said lysis buffer; and (h) freezing said cells semi-dry for storage. 25 61. A method for purifying a protein from a cell, comprising the steps of: (a) for each of a plurality of cell samples, lysing cells in each of said sample to produce a cell lysate, wherein said cell comprises a fusion protein having an affinity tag, and wherein said lysing step is performed using a paint 30 shaker: (b) separating each said lysate into a soluble fraction and a non-soluble fraction; (c) transferring each said soluble fraction into a different site of a multi-site array, wherein said transferring step is performed using a wide-open tip;

(d) contacting each said soluble fraction with a binding agent such that a complex between said fusion protein and binding agent is formed; (e) isolating each said fusion protein from the complex; and (f) storing each said fusion protein in a buffer of high viscosity. 5 A method for identifying whether a signal is positive, comprising the steps 62. of: (a) determining foreground and background signals for each spot locally and determining net signals from the difference between said foreground and 10 background signals; (b) determining the lower quartile, median, and upper quartile values of a first and second net signal distribution; (c) subtracting a first median value from said first net signal distribution, and subtracting a second median value from said second net signal distribution to obtain a first and second subtracted value, respectively; 15 (d) dividing said first subtracted value by the difference between said upper and lower quartile values of said first signal distribution, and dividing said second scaled value by the difference between said upper and lower quartile values to obtain a first and second scaled value, respectively; 20 (e) determining a local median value of a scaled signal distribution of a neighborhood region, wherein said neighborhood region comprises a plurality of sites in the area; and (f) subtracting the local median value from the scaled signal to obtain a scaled excess value. 25 The method of Claim 62, wherein a positive signal indicates protein-probe 63. interaction. 64. The method of Claim 62, wherein the neighborhood region is two rows 30 above, two rows below, two columns to the left, and two columns to the right of the signal. The method of Claim 62, further comprising the step of excluding parallel 65. samples of scaled excess values if the difference between one of the sample 35

values and the average of the sample values is greater than three standard deviations of the error of the scaled excess value.

- 66. A method for identifying positive signals among signals measured with a plurality of different arrays, comprising the steps of:
 - (a) transforming signals measured with different arrays to generate transformed signals;
 - (b) correcting each said transformed signal by a method comprising subtracting from said transformed signal a local median signal to generate a corrected transformed signal, wherein said local median signal is the median of signals in a neighborhood region, said neighborhood region comprising one or more sites around site of said transformed signal; and
 - (c) comparing said each said corrected transformed signal to a threshold value, and identifying said corrected transformed signal as positive if said corrected transformed signal is greater than said threshold value;

wherein said array comprises a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support.

- 67. The method of claim 66, wherein said step of transforming comprises the steps of:
 - (a) determining for signals measured with each of said different arrays the lower quartile value, the median value, and the upper quartile value of the signal distribution;
 - (b) subtracting from signals measured with each of said different arrays said median value to obtain translated signals for said array; and
 - (c) dividing said translated signals by the difference between said upper and lower quartile values of said array, thereby generating said transformed signals.

68. The method of claim 66, wherein said neighborhood region consists of sites within an area two rows above, two rows below, two columns to the left, and two columns to the right of the transformed signal.

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69. The method of Claim 66, 67, or 68, wherein said positive signal indicates protein-probe interaction.

- The method of Claim 66, 67, or 68, further comprising the step of discarding data points, wherein said data points are measured at duplicate sites on the array; and wherein the variation between said duplicate sites is greater than three standard deviations.
- 71. The method of Claim 66, 67, or 68, further comprising the step of normalizing the corrected transformed signals using the formula:
 r + ε_r = G + ε_G / R + ε_R
 wherein G is a corrected transformed signal, R is a GST signal, ε_G is the error of G, ε_R is the error of R, and ε_r is the error of r.
- The method of Claim 20, wherein the multi-site array is a 96-site array.

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- 73. A method for making a positionally addressable array comprising the step of attaching a plurality of fusion proteins to a surface of a solid support, with each fusion protein being at a different position on the solid support, wherein the fusion protein comprises a first tag, a second tag, and a protein sequence encoded by genomic nucleic acid of an organism.
- 74. The method of Claim 62 or 65, further comprising the steps of:

 (a) averaging the values of the signals of two duplicate spots to obtain an average value; and

 (b) determining whether said average value is greater than three standard deviations of the error of the scaled excess value, wherein a signal of said spot is positive if said average value is greater than three standard deviations of the error of the scaled excess value.
 - 75. A method for determining the presence or absence of an enzymatic activity in a protein comprising the steps of:(a) contacting a probe that is a substrate for said enzymatic activity with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support; and

(b) detecting any catalysis of said substrate at a position on the solid support; wherein said catalysis at a position on the solid support indicates that the protein at said position has said enzymatic activity.

- 5 76. A method for determining the presence or absence of an enzyme substrate in a protein comprising the steps of:
 - (a) contacting a probe that is an enzyme for said enzyme substrate with a positionally addressable array comprising a plurality of proteins, with each protein being at a different position on a solid support; and
 - (b) detecting any catalysis of said substrate at a position on the solid support; wherein said catalysis at a position on the solid support indicates that the protein comprises said enzyme substrate.
 - 77. The array of Claim 1, 3, 4, or 5, wherein the proteins are attached to the solid support via a biotin tag.
 - 78. The method of Claim 15, wherein the first tag is found at the carboxy-terminal end of the protein and the second tag is found at the amino-terminal end of the protein.
 - 79. The method of Claim 22, wherein the affinity tag is biotin.
 - 80. The array of Claim 1, 3, 4, or 5, wherein the solid support comprises a nitrocellulose-coated slide.

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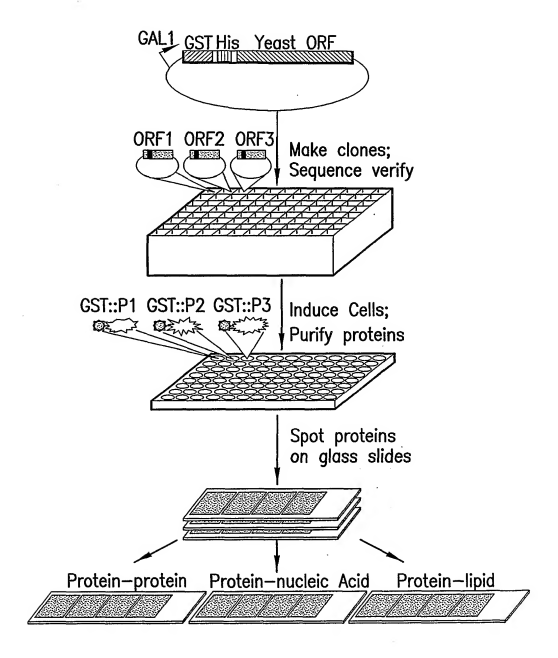


FIG. 1A

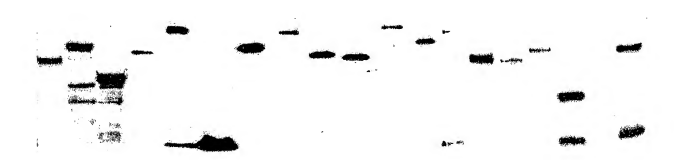


FIG.1B

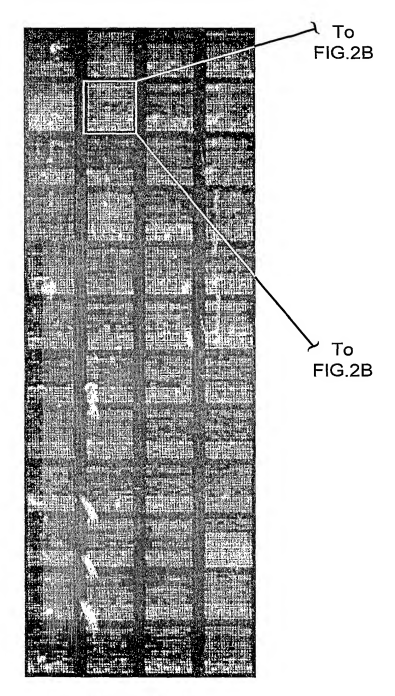
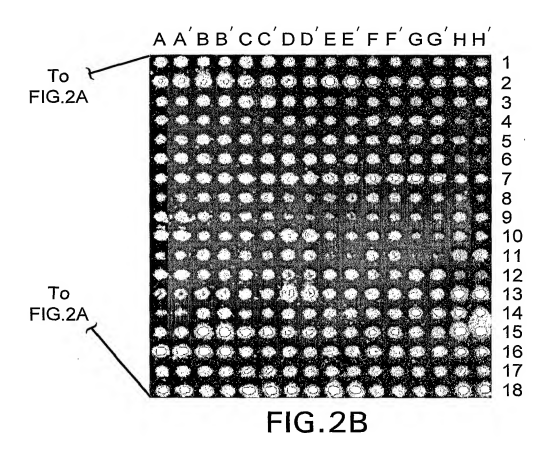


FIG.2A



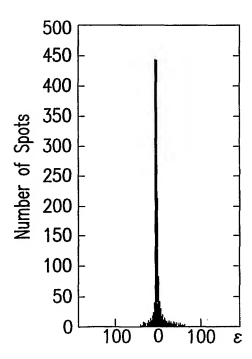


FIG.2C

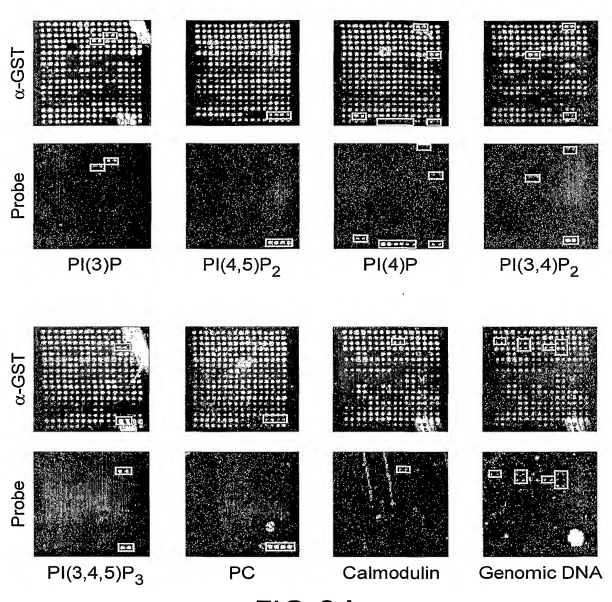


FIG.3A

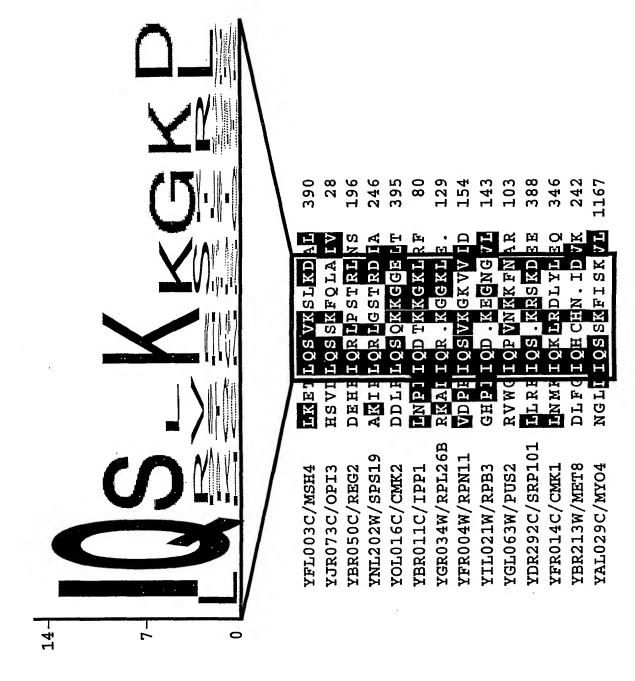


FIG. 5B

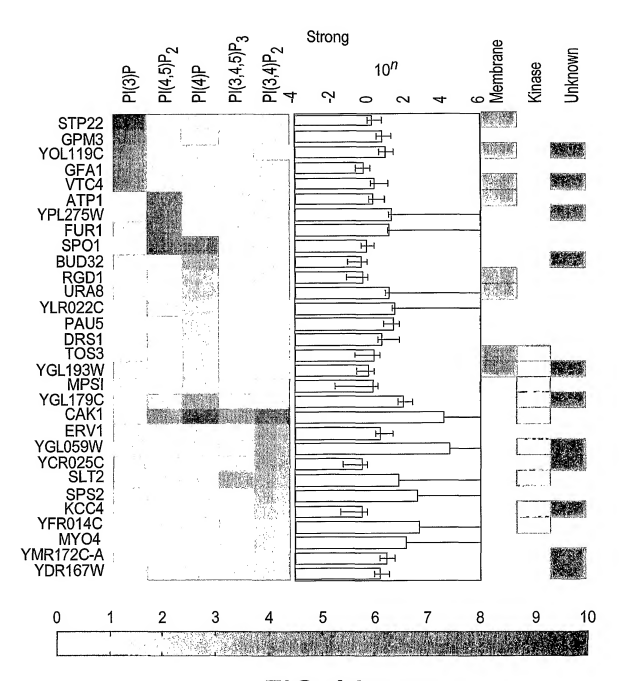


FIG.4A



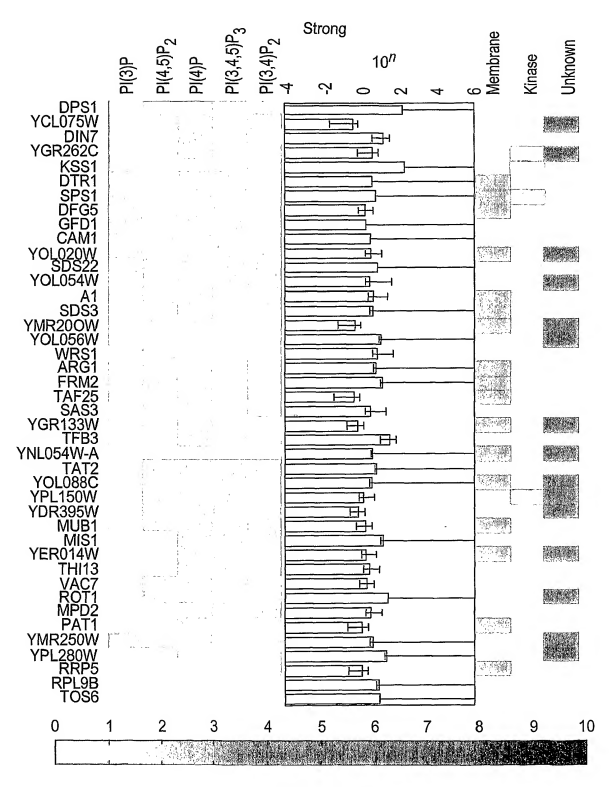


FIG.4B

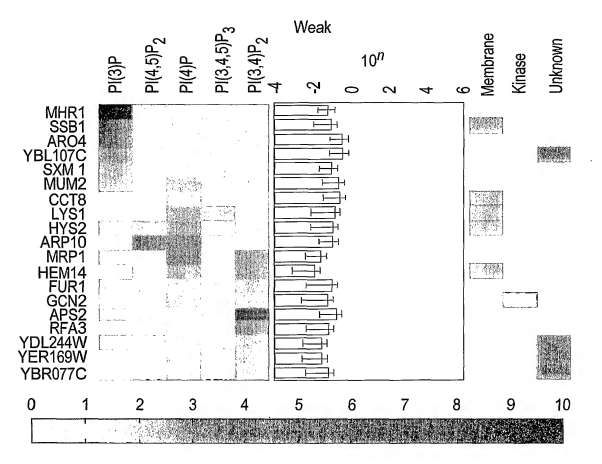
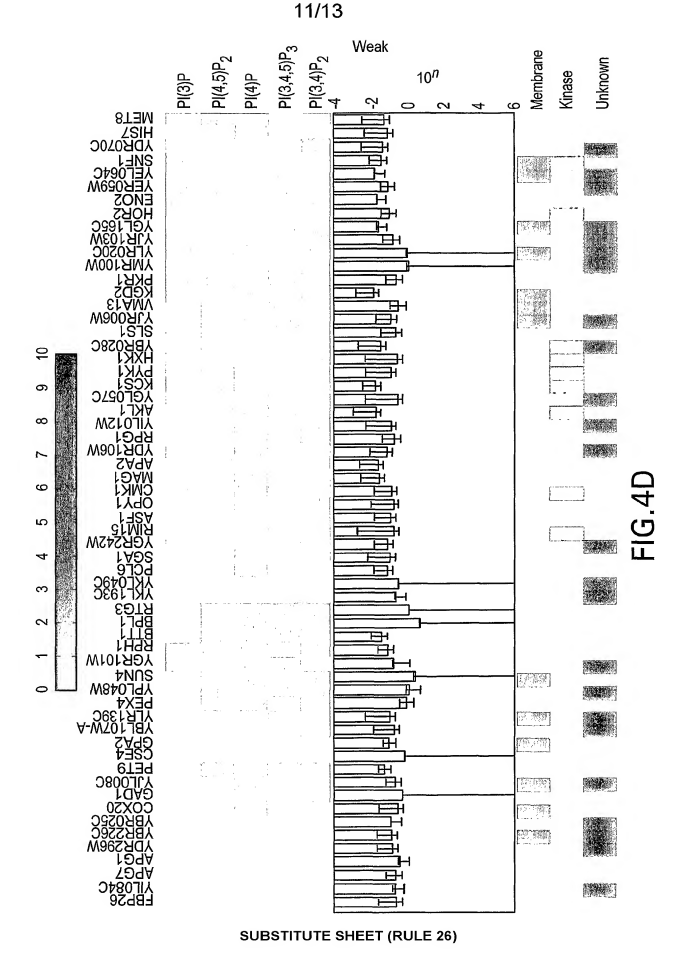
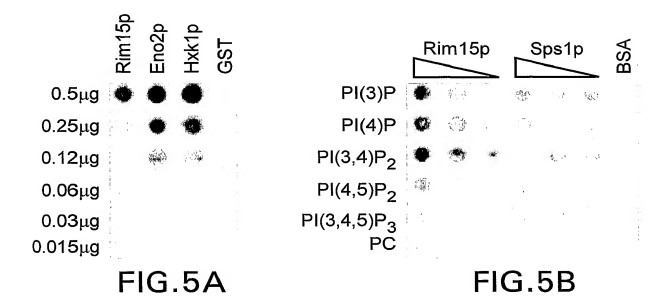
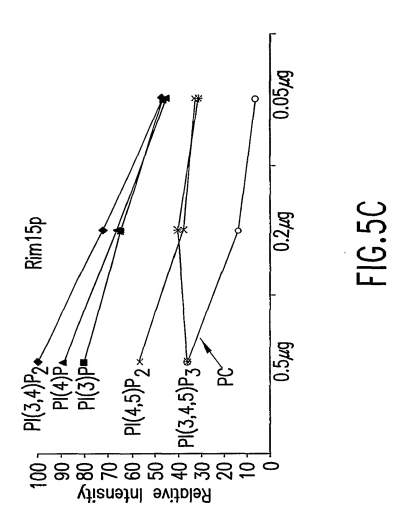


FIG.4C







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International application No. PCT/US02/14982

A. CLASSIFICATION OF SUBJECT MATTER				
` '	:A61K 38/00 :530/300			
	to International Patent Classification (IPC) or to both national classification and IPC			
B. FIEI	LDS SEARCHED			
Minimum d	locumentation searched (classification system followed by classification symbols)			
U.S. :	422/50,68.1; 435/7.1,7.2;530/300,333,350; 514/2			
Documentat	tion searched other than minimum documentation to the extent that such documents are	included in the fields		
searched				
Electronic o	data base consulted during the international search (name of data base and, where practicab	le, search terms used)		
Please Sec	e Extra Sheet.			
	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
Y	US 6,064,754 A (PAREKH et al.) 16 May 2000, see entire	1-57, 73,		
	document.	75-80		
Y	US 5,965,124 A (FEINBERG et al.) 12 October 1999, see entire	1-57, 73,		
	document.	75-80		
Y	US 6,146,830 A (FRIEND et al.) 14 November 2000, see entire	58 and 59		
	document.			
Y	US 6,197,599 A (CHIN et al.) 06 March 2001, see entire document.	58 and 59		
	·			
3.7	710 5 0 5 5 0 6 1 (COYIOTHIA 1) 00 T 1			
Y	US 5,866,362 A (COUSENS et al.) 02 February 1999, see entire	60		
	document.			
X Furtl	her documents are listed in the continuation of Box C See patent family annex.			
* Sp	ecial categories of cited documents: "I" later document published after the in date and not in conflict with the ap			
	oument defining the general state of the art which is not considered the principle or theory underlying to be of particular relevance			
	elier document published on or after the international filing date. "X" document of particular relevance;			
	considered novel or cannot be consid	юнея то шлогля ян шлентия втер		
	ed to establish the publication date of another citation or other solal reason (as specified) "Y" document of particular relevance;			
	considered to involve an inventive sk cument referring to an oral disclosure, use, exhibition or other with one or more other such does sans obvious to a person skilled in the a	ments, such combination being		
"P" do	cument published prior to the international filing date but later "&" document member of the same pater			
Date of the actual completion of the international search Date of mailing of the international search report				
09 AUGUST 2002				
Name and mailing address of the ISA/US Authorized officer				
Commissioner of Patents and Trademarks Box PCT ARDIV MARSCHEE ARDIV MARSCHEE				
Washington, D.C. 20231				
Facsimile N	lo. (703) 305-3230 Telephone No. (703) 308-0196	~		

International application No.
PCT/US02/14982

C (Continua	ation). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 5,846,819 A (PAUSCH et al.) 08 December 1998, see entire document.	60
Y	US 5,262,322 A (LIU et al.) 16 November 1993, see entire document.	60
Y	US 5,965,389 A (RAYMOND et al.) 12 October 1999, see entire document.	60
X 	US 5,720,928 A (SCHWARTZ) 24 February 1998, see entire document.	62
Y	· · · · · · · · · · · · · · · · · · ·	63-72, 74
Y	US 6,122,408 A (FANG et al.) 19 September 2000, see entire document.	62-72,74
Y	US 6,061,476 A (NICHANI) 09 May 2000, see entire document.	62-72,74
Y	US 6,075,875 A (GU) 13 June 2000, see entire document.	62-72, 74

International application No. PCT/US02/14982

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:		
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	
2.	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:	
s	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	
This Into	ernational Searching Authority found multiple inventions in this international application, as follows:	
Pl	ease See Extra Sheet.	
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.	
2. X	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.	
s	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:	
4.	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	
Remark	on Protest	
	No protest accompanied the payment of additional search fees.	

International application No. PCT/US02/14982

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

CAS, BOSIS, EMBASE, MEDLINE, WPI, BIOTECH ABS, covering search terms: protein, array, support, specie, yeast, culture, regulatory, expression, inducer, signal, positive, subtracted, foreground, background

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s) 1-19, 29-48, 50, 51, 73, 77, 78, and 80, drawn to a plurality of proteins on an array accounting for the majority of the proteins for a single specie.

Group II, claim(s) 20-24, 61, and 79, drawn to making protein samples via regulatory sequences and an inducer following by cell lysis.

Group III, claim(s) 25-28, 44-49, and 55-57, drawn to methods of detection of a lipid-binding protein via array usage.

Group IV, claims 52-54, drawn to methods of labeling and detecting biotinylated proteins.

Group V, claims 58 and 59, drawn to determination methods for posttranslational protein modification.

Group VI, claim 60, drawn to method of yeast culture preparation.

Group VII, claims 62-72 and 74, drawn to methods of identification of a positive signal via signal processing.

Group VIII, claims 75 and 76, drawn to methods of enzymatic activity determination.

The inventions listed as Groups I - VIII do not relate to a single inventive concept under PCT Rule 18.1 because, under PCT Rule 18.2, they lack the same or corresponding special technical features for the following reasons: Each of the Groups are directed to distinct and/or independent subject matter as listed for each group above. Group I is directed to protein arrays containing the majority of proteins for a single specie. Group II is directed to making proteins via regulatory expression controls and inducers. Group III is directed to detection of a particular lipid-binding protein which is not required in any other group. Group IV is directed to labeling and detecting biotinylated proteins which also is not required in any other group. Group V is directed to methods of detecting posttranslational protein modification. Group VI is directed to yeast culture preparation. Group VII is directed to methods of signal processing. Group VIII is directed to enzyme activity determination which is also not required in any other Group. Therefore, each Group is directed to a distinct and different Special Technical Feature which supports this Lack of Unity determination.